

**LIETUVOS SVEIKATOS MOKSLŲ UNIVERSITETO LIGONINĖ
KAUNO KLINIKOS**

(Adresatas (perkančioji organizacija))

PASIŪLYMAS**DĖL EKSPLOATACINIŲ PRIEDŲ ELEKTROCHIRURGINIAMS PRIETAISAMS PIRKIMO**

2025-01-20 Nr. _____

(Data)

Kaunas

(Sudarymo vieta)

1 lentelė

TIEKĖJO REKVIZITAI

Tiekėjo pavadinimas <i>//Jeigu dalyvauja ūkio subjektų grupė, surašomi visi dalyvių pavadinimai/</i>	UAB "Sorimpeksas"
Tiekėjo adresas <i>//Jeigu dalyvauja ūkio subjektų grupė, surašomi visi dalyvių adresai/</i>	Šiaulių 16a, 44353, Kaunas
Įmonės kodas, PVM mokėtojo kodas	135733248, LT357332412
Atsiskaitomosios sąskaitos numeris, bankas, banko kodas	LT647300010002276949 AB Swedbank, banko kodas 73000
Įmonės vadovo pareigos, vardas, pavardė	Direktorius Ramūnas Žalnerauskas
Už pasiūlymą atsakingo asmens vardas, pavardė	
Už sutarties vykdymą atsakingo asmense. pašto adresas, telefono numeris, el. pašto adresas	
Telefono numeris	8 37 361766
Fakso numeris	
El. pašto adresas	info@sorimpeksas.com

Šiuo pasiūlymu pažymime, kad sutinkame su visomis pirkimo sąlygomis, nustatytomis:

- 1) atviro konkurso skelbime, paskelbtame Viešųjų pirkimų įstatymo nustatyta tvarka;
- 2) kituose pirkimo dokumentuose (jų paaiškinimuose, papildymuose).

Pasirašydamas CVP IS priemonėmis pateiktą pasiūlymą saugiu elektroniniu parašu, patvirtinu, kad dokumentų skaitmeninės kopijos ir elektroninėmis priemonėmis pateikti duomenys yra tikri.

2 lentelė

SUBTIEKĖJO REKVIZITAI

Eil. Nr.	Subtiekėjo (-ų) pavadinimas (-ai), adresas (-ai)

*Pastaba: pildoma, jei tiekėjas ketina pasitelkti subtiekėją (-us)

3 lentelė

PASIŪLYMO KAINA

**Kainų pasiūlymas užpildytas pirkimo dokumentų 6 priede „Kainų pasiūlymo lentelė“
(dokumentas pateikiamas redaguojamu formatu)**

4 lentelė

PATEIKIAMŲ DOKUMENTŲ SĄRAŠAS

Eil.Nr.	Pateiktų dokumentų pavadinimas	Dokumen to puslapių skaičius	Failo, kuriame yra dokumentas, pavadinimas
1.	Konfidencialu. Direktorius igaliojimas (2024 E)	1	Konfidencialu. Direktorius igaliojimas (2024 E)
2.	Konfidencialu. CE sertifikatai	150	Konfidencialu. CE sertifikatai
3.	EBVPD pažyma	14	EBVPD pažyma
4.	7_7 priedas DEKLARACIJA DĖL TIEKĖJO ATSAKINGŲ ASMENŲ	1	7_7 priedas DEKLARACIJA DĖL TIEKĖJO ATSAKINGŲ ASMENŲ
5.	8_8 priedas Tiekėjo deklaracija	1	8_8 priedas Tiekėjo deklaracija
6.	4.1 pirkio dalis	1	4.1 pirkio dalis
7.	4.2 pirkio dalis	1	4.2 pirkio dalis
8.	5 pirkimo dalis	1	5 pirkimo dalis
9.	Bissinger	63	Bissinger
10.	E2100	1	E2100
11.	Kempinėlės	1	Kempinėlės
12.	Konfidencialu. 9.1 pirkimo dalis	1	Konfidencialu. 9.1 pirkimo dalis
13.	Tiekėjo patvirtinimas	1	Tiekėjo patvirtinimas

Pastaba. Perkančioji organizacija atmes tiekėjo pasiūlymą, kaip neatitinkantį pirkimo dokumentuose nustatytų reikalavimų, jeigu kartu su pasiūlymu nebus pateikti pirkimo sąlygų 5.11.2, 5.11.9, 5.11.11 punktuose nurodyti dokumentai.

Pasiūlymas galioja iki termino, nustatyto pirkimo dokumentuose.

Primintina, kad pasiūlyme nurodytos kainos bei įkainiai, taip pat nuolaidos dydis ar įkainio bazė, tiekėjo siūlomų prekių gamintojai, pavadinimai, modeliai, tiekėjo siūlomų prekių techninės specifikacijos, nurodomos užpildant perkančiosios organizacijos pateiktas lenteles, gaminio naudotojo instrukcija, tiekėjo siūlomų prekių atitiktį techninės specifikacijos reikalavimams įrodantys dokumentai - brošiūros, aprašymai, instrukcijos - nėra konfidenciali informacija (plačiau skaityti¹).

Pasiūlymo konfidencialią informaciją sudaro: Konfidencialu. Direktorius igaliojimas (2024 E); Konfidencialu. CE sertifikatai; Konfidencialu. 9.1 pirkimo dalis

(Tiekėjo arba jo įgalioto asmens
pareigų pavadinimas)

(Parašas)

Vardas, pavardė

¹ https://vpt.lrv.lt/uploads/vpt/documents/files/mp/konfidenciali_informacija.pdf

Tiekėjo pavadinimas (nurodyti): UAB Sorimpeksas

Pirkimo dalies Nr.	Pavadinimas	Modelis, katalogo numeris, gamintojo pavadinimas	Mato vnt.	Orientacinis kiekis	Vieneto kaina Eur (be PVM)	Kaina viso Eur (be PVM)	Kaina viso Eur (su PVM)
4.	Elektrochirurginio prietaiso priedai: bipoliariniai kabeliai						
4.1	Bipoliarinis kabelis	Bipolinis kabelis, Bissinger 80100033	vnt.	250	57,00	14250,00	17242,50
4.2	Bipoliarinis kabelis	Bipolinis kabelis, Bissinger 80100067	vnt.	60	55,00	3300,00	3993,00
Bendra 4 pirkimo dalies pasiūlymo kaina EUR (be PVM):							17550,00
PVM suma:							3685,50
Bendra 4 pirkimo dalies pasiūlymo kaina EUR (su PVM):							21235,50
5.	Elektrochirurginio prietaiso priedai: pasyvių vienkartinį elektrodą kabeliai						
5.1	Pasyvaus vienkartinio elektrodo kabelis	Neutralių elektrodų pajungimo kabelis, Bissinger 89101051	vnt.	40	66,00	2640,00	3194,40
5.2	Pasyvaus vienkartinio elektrodo kabelis	Neutralių elektrodų pajungimo kabelis, Bissinger 89101053	vnt.	40	66,00	2640,00	3194,40
Bendra 5 pirkimo dalies pasiūlymo kaina EUR (be PVM):							5280,00
PVM suma:							1108,80
Bendra 5 pirkimo dalies pasiūlymo kaina EUR (su PVM):							6388,80

Eksplotacinių priedų elektrochirurginiams prietaisams techninė specifikacija

Pirkimo dalies Nr.	Pavadinimas (specifikacija)	Reikalaujamos parametrų reikšmės	Gamintojo „BOWA“ kodas REF arba lygiavertis	Siūlomos parametrų reikšmės
4.	Elektrochirurginio prietaiso priedai: bipoliariniai kabeliai			
4.1	Bipoliarinis kabelis (orientacinis kiekis 250 vnt.)	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis $\geq 4,5$ m; 5. Su europinio tipo plokščia (arba lygiaverte) jungtimi bipoliariniams pincetams; 6. Su $28,5 \text{ mm} \pm 0,5 \text{ mm}$ pločio 2 kontaktų tarptautine jungtimi pajungimui prie generatoriaus; 	Bissinger 80100033	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis 5 m; 5. Su europinio tipo plokščia jungtimi bipoliariniams pincetams; 6. Su 29mm pločio 2 kontaktų tarptautine jungtimi pajungimui prie generatoriaus; <i>4.1 pirkimo dalis;</i> <i>Bissinger 53 psl.</i>
4.2	Bipoliarinis kabelis (orientacinis kiekis 60 vnt.)	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis $\geq 4,5$ m; 5. Su europinio tipo plokščia (arba lygiaverte) jungtimi bipoliariniams pincetams; 6. Su $22 \text{ mm} \pm 0,5 \text{ mm}$ pločio 2 kontaktų tarptautine jungtimi pajungimui prie generatoriaus; 	Bissinger 80100067	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis 5 m; 5. Su europinio tipo plokščia jungtimi bipoliariniams pincetams; 6. Su 22mm pločio 2 kontaktų tarptautine jungtimi pajungimui prie generatoriaus; <i>4.1 pirkimo dalis;</i> <i>Bissinger 53 psl.</i>
5.	Elektrochirurginio prietaiso priedai: pasyvių vienkartinį elektrodų kabeliai			
5.1	Pasyvaus vienkartinio elektrodo kabelis (orientacinis kiekis 40 vnt.)	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis $\geq 4,5$ m; 5. Prijungimas – 2 kontaktų REM tarptautinis. 	Bissinger 89101051	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis 4,5 m; 5. Prijungimas – 2 kontaktų REM tarptautinis. <i>5 pirkimo dalis;</i> <i>Bissinger 55 psl.</i>
5.2	Pasyvaus vienkartinio elektrodo kabelis (orientacinis kiekis 40 vnt.)	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis $\geq 4,5$ m; 5. Prijungimas – 2 kontaktų NON REM tarptautinis. 	Bissinger 89101053	<ol style="list-style-type: none"> 1. Sterilizuojamas; 2. Daugkartinio naudojimo; 3. Su papildomomis apsaugomis nuo nulankstymo prie jungčių; 4. Kabelio ilgis 4,5 m; 5. Prijungimas – 2 kontaktų NON REM tarptautinis <i>5 pirkimo dalis;</i> <i>Bissinger 55 psl.</i>

Pastabos, papildomi reikalavimai:

1. Lentelėje pateikti firmų pavadinimai ir kataloginiai numeriai jokios komercinės reikšmės neturi, o tik nurodo technines gaminių charakteristikas aprašantį informacijos šaltinį. Gali būti siūlomos nurodytų gamintojų prekės arba joms lygiaverčiai, ne blogesnių techninių ir saugumo charakteristikų, kitų firmų gaminiai.
2. Visi priedai techniškai suderinami su LSMU ligoninėje Kauno klinikose naudojamais elektrochirurginiais prietaisais BOWA ARC 350.
3. Garantinis terminas ne mažiau kaip 12 mėnesių (reikalavimas taikomas daugkartinio naudojimo priedams).
4. Pasiūlymo priede turi būti pateikti katalogai, prospektai ar kita informacija su siūlomų prekių iliustracijomis.
5. Siūlomos prekės turi būti ženklinamos CE ženklu (*kartu su pasiūlymu konkursui privaloma pateikti CE sertifikato arba EB atitikties deklaracijos kopiją*).

6. Viešojo pirkimo komisijai pareikalavus, išbandymui turi būti pateikti siūlomų prekių pavyzdžiai.
7. Prekių pristatymo išlaidos įskaičiuotos į pasiūlymo kainą.

Europos bendrasis viešųjų pirkimų dokumentas (EBVPD)

I dalis. Informacija apie pirkimo procedūrą ir perkančiąją organizaciją ar perkantįjį subjektą

Informacija apie paskelbimą

Skelbimo numeris OL S (tik tarptautiniams pirkimams):

-

Skelbimo numeris CVP IS (kur rasti?)

-

Perkančiosios organizacijos / Perkančiojo subjekto tapatybė

Oficialus pavadinimas:

LSMUL KAUNO KLINIKOS

Šalis:

Lietuva

Informacija apie pirkimo procedūrą

Procedūros tipas

Atvira

Pavadinimas:

EKSPLOATACINIAI PRIEDAI ELEKTROCHIRURGINIAMS PRIETAISAMS

Trumpas aprašymas:

EKSPLOATACINIAI PRIEDAI ELEKTROCHIRURGINIAMS PRIETAISAMS

Perkančiosios organizacijos ar perkančiojo subjekto (jei taikoma) priskirtas dokumento numeris:

-

II dalis. Informacija apie ekonominės veiklos vykdytoją

A. Informacija apie ekonominės veiklos vykdytoją

Tiekėjo pavadinimas arba vardas ir pavardė (jei fizinis asmuo):

UAB "Sorimpeksas"

Gatvė ir namo numeris:

Šiaulių 16a.

Pašto kodas:

LT44353

Miestas:

Kaunas

Šalis:

Lietuva

Interneto adresas (jei yra):

-

E. paštas:

[redacted]

Telefonas:

[redacted]

Asmuo ar asmenys ryšiams:

[redacted]

PVM mokėtojo kodas, jei yra:

LT357332412

Jei PVM mokėtojo kodo nėra, nurodykite kitą nacionalinį identifikacinį numerį (Lietuvoje - įmonės kodą)

-

Ar ekonominės veiklos vykdytojas yra labai maža, mažoji ar vidutinė įmonė?

☒ Taip

☐ Ne

Tik tuo atveju, kai pirkimas rezervuotas: ar ekonominės veiklos vykdytojas yra globojama darbo grupė (neįgalųjų socialinė įmonė), socialinė įmonė? Ar jis vykdys sutartį pagal globojamų darbo grupių (neįgalųjų socialinių įmonių) užimtumo programas?

☐ Taip

☒ Ne

Jei taikoma, ar ekonominės veiklos vykdytojas įtrauktas į oficialų patvirtintų ekonominės veiklos vykdytojų sąrašą arba ar jis turi lygiavertį sertifikatą (pvz., pagal nacionalinę (išankstinę) kvalifikacijos vertinimo sistemą)? Lietuvos tiekėjai renkasi „ne“

☐ Taip

☒ Ne

- Be to, užpildykite trūkstamą informaciją IV dalies A, B, C arba D skirsniuose, atsižvelgdami į konkretų atvejį TIK jei to reikalaujama atitinkamame skelbime arba pirkimo dokumentuose:

e) Ar ekonominės veiklos vykdytojas galės pateikti sertifikatą dėl socialinio draudimo įmokų ir mokesčių mokėjimo arba pateikti informaciją, kuri leistų perkančiajai organizacijai ar perkančiajam subjektui jį gauti tiesiogiai naudojantis prieiga prie bet kurios iš valstybių narių nemokamos nacionalinės duomenų bazės?

☒ Taip

☐ Ne

Jei atitinkami dokumentai prieinami elektroniniu būdu, nurodykite:

https://draudejai.sodra.lt/draudeju_viesi_duomenys/

Ar ekonominės veiklos vykdytojas pirkimo procedūroje dalyvauja kartu su kitais? Žymima TAIP, jei pasiūlymą teikia ūkio subjektų grupė (konsorciumas) pagal jungtinės veiklos sutartį

☐ Taip

☒ Ne

Jei pirkimas padalintas į dalis, nuoroda į pirkimo dalį (-is), dėl kurios (-ių) ekonominės veiklos vykdytojas nori dalyvauti konkurse:

4, 5, 7, 8, 9 pirkimo dalys

B. Informacija apie ekonominės veiklos vykdytojo teisinius atstovus #1

- Šis skirsnis pildomas, jeigu tiekėjo vadovas įgalioja kitą asmenį pasirašyti pasiūlymą, bendrauti su pirkimo vykdytoju, įgalioja atstovauti ir pasirašyti EBVPD, bendrauti su pirkimo vykdytoju dėl EBVPD pateiktos informacijos, teikiamų kvalifikaciją ir pašalinimo pagrindų nebuvimą pagrindžiančių dokumentų, dėl pasiūlymo ir pan.

Jei taikytina, nurodykite asmens (-ų), įgalioto (-ų) atstovauti ekonominės veiklos vykdytojui šios pirkimo procedūros tikslais, vardą ir pavardę ir adresą:

Vardas



Pavardė

Gimimo data

-

Gimimo vieta

-

Gatvė ir namo numeris:

Šiaulių 16 a.

Pašto kodas:

LT44353

Miestas:

Kaunas

Šalis:

Lietuva

E. paštas:

-

Telefonas:

-

Pareigos arba statusas:

-

Prireikus pateikite išsamią informaciją apie atstovavimą (formą, aprėptį, paskirtį ir t. t.):

-

C. Informacija apie rėmimąsi kitų subjektų pajėgumais

Ar siekdamas patenkinti IV dalyje nurodytus atrankos kriterijus ir V dalyje nurodytus kriterijus bei taisykles (jei tokių yra) ekonominės veiklos vykdytojas remiasi kitų subjektų pajėgumais?

☐ Taip

☒ Ne

D. Informacija apie subrangovus, kurių pajėgumais ekonominės veiklos vykdytojas nesiremia

- (Skirsnį reikia pildyti, tik jei šios informacijos aiškiai reikalauja perkančioji organizacija ar perkantysis subjektas.)

Ar ekonominės veiklos vykdytojas ketina kurias nors sutarties dalis subrangos sutartimi pavesti atlikti trečiosioms šalims?

☐ Taip

☒ Ne

- Jei perkančioji organizacija ar perkantysis subjektas aiškiai prašo šios informacijos, šalia informacijos pagal šį skirsnį, pateikite pagal šios dalies A ir B skirsnius ir III dalį reikalaujamą informaciją apie kiekvieną susijusį subrangovą (subrangovų kategorijas).

III dalis. Pašalinimo pagrindai

A. Su baudžiamaisiais nuosprendžiais susiję pagrindai

Direktyvos 2014/24/ES 57 straipsnio 1 dalyje nustatyti šie pašalinimo pagrindai

A1. Dalyvavimas nusikalstamos organizacijos veikloje (VPĮ 46 str. 1 d. 1 p.)

Ar pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo nuteistas galutiniu teismo sprendimu už dalyvavimą nusikalstamos organizacijos veikloje, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia? Kaip apibrėžta 2008 m. spalio 24 d. Tarybos pamatinio sprendimo 2008/841/TVR dėl kovos su organizuotu nusikalstamumu 2 straipsnyje (OL L 300, 2008 11 11, p. 42).

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

A2. Korupcija (VPĮ 46 str. 1 d. 2 p.)

Ar pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo nuteistas galutiniu teismo sprendimu už korupciją, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo

laikotarpis tebesitęsia? Kaip apibrėžta Konvencijos dėl kovos su korupcija, susijusia su Europos Bendrijų pareigūnais ar Europos Sąjungos valstybių narių pareigūnais, 3 straipsnyje (OL C 195, 1997 6 25, p. 1) ir 2003 m. liepos 22 d. Tarybos pamatinio sprendimo 2003/568/TVR dėl kovos su korupcija privačiame sektoriuje 2 straipsnio 1 dalyje (OL L 192, 2003 7 31, p. 54). Į pašalinimo pagrindus taip pat įtraukta korupcija, kaip apibrėžta perkančiosios organizacijos (perkančiojo subjekto) arba ekonominės veiklos vykdytojo nacionalinėje teisėje.

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

A3. Sukčiavimas (VPĮ 46 str. 1 d. 3 p.)

Ar pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo nuteistas galutiniu teismo sprendimu už sukčiavimą, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia? Pagal Europos Bendrijų finansinių interesų apsaugos konvencijos 1 straipsnį (OL C 316, 1995 11 27, p. 48).

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

A4. Teroristiniai nusikaltimai arba su teroristine veikla susiję nusikaltimai (VPĮ 46 str. 1 d. 5 p.)

Ar pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo nuteistas galutiniu teismo sprendimu už teroristinius nusikaltimus arba

su teroristine veikla susijusius nusikaltimus, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia? Kaip apibrėžta 2002 m. birželio 13 d. Tarybos pamatinio sprendimo dėl kovos su terorizmu 1 ir 3 straipsniuose (OL L 164, 2002 6 22, p. 3). Į pašalinimo pagrindus taip pat įtrauktas nusikalstamos veikos kurstymas, pagalba ar bendrininkavimas ją vykdant arba kėsiniimasis ją įvykdyti, kaip nurodyta to pamatinio sprendimo 4 straipsnyje.

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

A5. Pinigų plovimas arba teroristų finansavimas (VPĮ 46 str. 1 d. 6 p.)

Ar pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo nuteistas galutiniu teismo sprendimu už pinigų plovimą arba teroristų finansavimą, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia? Kaip apibrėžta 2005 m. spalio 26 d. Europos Parlamento ir Tarybos direktyvos 2005/60/EB dėl finansų sistemos apsaugos nuo jos panaudojimo pinigų plovimui ir teroristų finansavimui 1 straipsnyje (OL L 309, 2005 11 25, p. 15).

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

A6. Vaikų darbas ir kitos prekybos žmonėmis formos (VPĮ 46 str. 1 d. 7 p.)

Ar pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo,

sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo nuteistas galutiniu teismo sprendimu už vaikų darbą arba kitas prekybos žmonėmis formas, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia? Kaip apibrėžta 2011 m. balandžio 5 d. Europos Parlamento ir Tarybos direktyvos 2011/36/ES dėl prekybos žmonėmis prevencijos, kovos su ja ir aukų apsaugos, pakeičiančios Tarybos pamatinį sprendimą 2002/629/TVR, 2 straipsnyje (OL L 101, 2011 4 15, p. 1).

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

B. Su mokesčių ar socialinio draudimo įmokų mokėjimu susiję pagrindai **Direktyvos 2014/24/ES 57 straipsnio 2 dalyje nustatytos šios pašalinimo priežastys**

B1. Mokesčių mokėjimas VPĮ 46 str. 3 d.

Ar ekonominės veiklos vykdytojas pažeidė savo pareigas, susijusias su mokesčių mokėjimu, tiek šalyje, kurioje yra įsisteigęs, tiek perkančiosios organizacijos ar perkančiojo subjekto valstybėje narėje, jei tai nėra jo įsisteigimo šalis?

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

B2. Socialinio draudimo įmokų mokėjimas VPĮ 46 str. 3 d.

Ar ekonominės veiklos vykdytojas pažeidė savo pareigas, susijusias su socialinio draudimo įmokų mokėjimu, tiek šalyje, kurioje yra įsisteigęs, tiek perkančiosios organizacijos ar perkančiojo subjekto valstybėje narėje, jei tai nėra jo įsisteigimo šalis?

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☒ Taip

☐ Ne

URL

https://draudejai.sodra.lt/draudeju_viesi_duomenys/

Kodas

191630223

Emitentas

Valstybinio socialinio draudimo fondo valdyba prie Socialinės apsaugos ir darbo ministerijos

C. Su nemokumu, interesų konfliktu ar profesiniais nusižengimais susiję pagrindai

Direktyvos 2014/24/ES 57 straipsnio 4 dalyje nustatyti šie pašalinimo pagrindai

C10. Su kitais ekonominės veiklos vykdytojais sudaryti susitarimai, kuriais siekta iškreipti konkurenciją (VPĮ 46 str. 4 d. 1 p.)

Ar ekonominės veiklos vykdytojas su kitais ekonominės veiklos vykdytojais yra sudaręs susitarimų, kuriais siekta iškreipti konkurenciją atliekamame pirkime?

Jūsų atsakymas

☐ Taip

☒ Ne

C11. Rimti profesiniai pažeidimai VPĮ 46 str. 4 d. 7 p., VPĮ 46 str. 6 d. 3 p.

Pirkimams pradėtiems nuo 2022-01-01: Ar ekonominės veiklos vykdytojas yra padaręs rimtą profesinį pažeidimą, kaip nurodyta žemiau?:

a) yra padaręs finansinės atskaitomybės ir audito teisės aktų pažeidimą ir nuo jo padarymo dienos praėjo mažiau kaip vieni metai; **Nuo 2022-08-12**

pildydamas EBVPD tiekėjas yra informuotas ir supranta, kad finansinės atskaitomybės ir audito teisės aktų pažeidimu taip pat gali būti laikomi atvejai, kai tiekėjas nepateikia privalomų finansinės atskaitomybės dokumentų Registrų centrui. Išsamiau: <https://vpt.lrv.lt/lt/naujienos/>

finansiniu-ataskaitu-nepateikimas-gali-tapti-kliutimi-dalyvauti-viesuosiuose-pirkimuose

b) neatitinka minimalių patikimo mokesčių mokėtojo kriterijų, nustatytų Lietuvos Respublikos mokesčių administravimo įstatymo 40¹ straipsnio 1 dalyje. Taikant šį tiekėjo pašalinimo iš pirkimo procedūros pagrindą, vadovaujamosi Lietuvos Respublikos mokesčių administravimo įstatymo 40¹ straipsnio 1 dalyje nustatytais terminais, juos skaičiuojant nuo Mokesčių administravimo įstatymo 40¹ straipsnio 1 dalyje nurodytų pažeidimų padarymo dienos, tačiau visais atvejais šie terminai negali būti ilgesni negu 3 metai;

c) yra padaręs draudimo sudaryti draudžiamus susitarimus, įtvirtinto Lietuvos Respublikos konkurencijos įstatyme ar panašaus pobūdžio kitos valstybės teisės akte, pažeidimą ir nuo jo padarymo dienos praėjo mažiau kaip 3 metai;

d) yra padaręs bet kokį kitą rimtą profesinį pažeidimą, nenurodytą aukščiau, nuo kurio padarymo dienos praėjo mažiau kaip vieni metai?

Pirkimams pradėtiems iki 2022-01-01: Ar ekonominės veiklos vykdytojas yra pripažintas kaltu dėl sunkaus profesinio nusižengimo kaip nurodyta žemiau?

I. ar ekonominės veiklos vykdytojas yra padaręs profesinį pažeidimą, kai už finansinės atskaitomybės ir audito teisės aktų pažeidimus ekonominės veiklos vykdytojui ar jo vadovui paskirta administracinė nuobauda ar ekonominė sankcija, nustatytos Lietuvos Respublikos įstatymuose ar kitų valstybių teisės aktuose, ir nuo sprendimo, kuriuo buvo paskirta ši sankcija, įsiteisėjimo dienos arba nuo dienos, kai asmuo įvykdė administracinį nurodymą, praėjo mažiau kaip vieni metai?

II. Ar ekonominės veiklos vykdytojas yra padaręs kurį nors vieną iš žemiau nurodytų rimtų profesinių pažeidimų(taikoma tik tada kai, ir tik tiek, kiek apibrėžta kituose pirkimo dokumentuose):

a) profesinės etikos pažeidimas, kai nuo ekonominės veiklos vykdytojo pripažinimo nesilaikančiu profesinės etikos normų momento praėjo mažiau kaip vieni metai;

b) konkurencijos, darbuotojų saugos ir sveikatos, informacijos apsaugos, intelektinės nuosavybės apsaugos pažeidimas, už kurį ekonominės veiklos vykdytojui ar jo vadovui yra paskirta administracinė nuobauda ar ekonominė sankcija, nustatytos Lietuvos Respublikos ar kitų valstybių įstatymuose, kai nuo sprendimo, kuriuo buvo paskirta ši sankcija, arba nuo dienos, kai asmuo įvykdė administracinį nurodymą, įsiteisėjimo dienos praėjo mažiau kaip vieni metai;

c) draudimo sudaryti draudžiamus susitarimus, įtvirtinto Lietuvos Respublikos konkurencijos įstatyme ar panašaus pobūdžio kitos valstybės teisės akte, pažeidimas, kai nuo sprendimo paskirti Konkurencijos įstatyme ar kitos valstybės teisės akte nustatytą ekonominę sankciją įsiteisėjimo dienos praėjo mažiau kaip 3 metai;

d) ekonominės veiklos vykdytojas, kuris yra fizinis asmuo, arba ekonominės veiklos vykdytojo, kuris yra juridinis asmuo, kita organizacija ar jos padalinys, vadovas, kitas valdymo ar priežiūros organo narys ar kitas asmuo, turintis (turintys) teisę atstovauti ekonominės veiklos vykdytojui ar jį kontroliuoti, jo vardu priimti sprendimą, sudaryti sandorį, arba dalyvis, turintis balsų daugumą juridinio asmens dalyvių susirinkime, yra pripažintas kaltu dėl tyčinio bankroto, kaip jis apibrėžtas Lietuvos Respublikos įmonių bankroto įstatyme ar panašaus pobūdžio kitų valstybių teisės aktuose, kai nuo teismo sprendimo įsiteisėjimo dienos praėjo mažiau kaip 3 metai?

Jūsų atsakymas

☐ Taip

☒ Ne

C12. Interesų konfliktas dėl dalyvavimo pirkimo procedūroje (VPĮ 46 str. 4 d. 2 p.)

Ar ekonominės veiklos vykdytojas žino apie kokius nors [interesų konfliktus](#), kaip nurodyta nacionalinėje teisėje, atitinkamame skelbime ar pirkimo dokumentuose, kylančius dėl jo dalyvavimo pirkimo procedūroje?

Jūsų atsakymas

☐ Taip

☒ Ne

C13. Tiesioginis arba netiesioginis dalyvavimas rengiant šią pirkimo procedūrą (46 str. 4 d. 3 p.)

Ar ekonominės veiklos vykdytojas arba su juo susijusi įmonė konsultavo perkančiąją organizaciją ar perkantįjį subjektą arba kitaip dalyvavo rengiant pirkimo procedūrą?

Jūsų atsakymas

☐ Taip

☒ Ne

C14. Sutarties nutraukimas anksčiau laiko, žala ar kitos panašios sankcijos (VPĮ 46 str. 4 d. 6 p.)

Ar ekonominės veiklos vykdytojas turėjo tokios patirties: ankstesnė viešoji sutartis, ankstesnė sutartis su perkančiuoju subjektu arba ankstesnė koncesijos sutartis buvo nutraukta anksčiau laiko; arba buvo pareikalauta atlyginti su ankstesne sutartimi susijusią žalą ar skirtos kitos panašios sankcijos?

Lietuvoje (be kita ko) - ar ekonominės veiklos vykdytojas yra įtrauktas į nepatikimų tiekėjų sąrašą ?

Jūsų atsakymas

☐ Taip

●Ne

C15. Pripažinimas kaltu dėl faktų iškreipymo, informacijos nuslėpimo, negalėjimas pateikti reikalaujamų dokumentų ir su šia procedūra susijusios konfidencialios informacijos gavimas (46 str. 4 d. 4 p. ir 46 str. 4 d. 5 p.)

Ar ekonominės veiklos vykdytojas yra susijęs su vienu iš šių atvejų, kai jis :

- a) buvo labai iškreipęs faktus pateikdamas informaciją (**pateikęs melagingą informaciją**), reikalingą patikrinti, ar nėra pagrindų pašalinti, arba patikrinti atitiktį atrankos kriterijams;
- b) slėpė tokią informaciją;
- c) delsė pateikti patvirtinamuosius dokumentus, kurių reikalavo perkančioji organizacija ar perkantysis subjektas,
- d) siekė daryti neteisėtą įtaką perkančiosios organizacijos ar perkančiojo subjekto sprendimų priėmimo procesui, kad gautų konfidencialios informacijos, dėl kurios per pirkimo procedūrą įgytų nepagrįstą pranašumą, arba tyčia teikti klaidinančios informacijos, kuri gali turėti esminės įtakos sprendimams dėl pašalinimo, atrankos ar sutarties skyrimo?

Jūsų atsakymas

○Taip

●Ne

D. Išimtinai nacionaliniai pašalinimo pagrindai

Išimtinai nacionaliniai pašalinimo pagrindai, nurodyti atitinkamame skelbime ar pirkimo dokumentuose.

D1. Išimtinai nacionaliniai pašalinimo pagrindai (VPĮ 46 str. 1 d. 4 p.)

Pirkimams pradėtiems nuo 2022-01-01:

pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo **nuteistas galutiniu teismo sprendimu už nusikalstamą bankrotą**, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia?

Pirkimams pradėtiems iki 2022-01-01:

Ar ekonominės veiklos vykdytojas yra susijęs su vienu iš šių atvejų, kai:

- a) jis **neatitinka minimalių patikimo mokesčių mokėtojo kriterijų**, nustatytų Lietuvos Respublikos mokesčių administravimo įstatymo 40¹ straipsnio 1 dalyje ir dėl to laikomas padariusiu šiurkštų profesinį pažeidimą.

b) pats ekonominės veiklos vykdytojas ar bet kuris asmuo, kuris yra jo administracijos, valdymo ar priežiūros organo narys arba turi atstovavimo, sprendimo ar kontrolės įgaliojimus to ekonominės veiklos vykdytojo atžvilgiu, buvo **nuteistas galutiniu teismo sprendimu už nusikalstamą bankrotą**, o nuosprendis priimtas prieš ne daugiau kaip penkerius metus arba kai nuosprendyje aiškiai nustatytas pašalinimo laikotarpis tebesitęsia?

Jūsų atsakymas

☐ Taip

☒ Ne

Ar ši informacija ES valstybės narės duomenų bazėje nemokamai prieinama valdžios institucijoms?

☐ Taip

☒ Ne

IV dalis. Atrankos kriterijai

α. Visų atrankos kriterijų bendra nuoroda

Dėl atrankos kriterijų ekonominės veiklos vykdytojas pareiškia, kad jis atitinka visus reikalaujamus atrankos kriterijus

Jūsų atsakymas

☒ Taip

☐ Ne

Baigti

IV dalis. Baigiamieji pareiškimai

Ekonominės veiklos vykdytojai oficialiai pareiškia, kad II–V dalyse pateikta informacija yra tiksli ir teisinga ir kad ji pateikta visiškai suvokiant didelio faktų iškreipimo padarinius.

Ekonominės veiklos vykdytojai oficialiai pareiškia, kad pareikalavus gali nedelsdami pateikti nurodytus sertifikatus ir kitų formų įrodomuosius dokumentus, išskyrus tuos atvejus, kai:

a) perkančioji organizacija ar perkantysis subjektas turi galimybę atitinkamus patvirtinamuosius dokumentus tiesiogiai gauti naudodamiesi prieiga prie bet kurios iš valstybių narių nemokamos nacionalinės duomenų bazės (su sąlyga, kad ekonominės veiklos vykdytojas pateikė reikalingą informaciją (interneto adresą, išduodančiąją instituciją ar įstaigą, tikslias dokumentų nuorodas),

kuri perkančiajai organizacijai ar perkančiajam subjektui leidžia tai padaryti (pareikalavus dėl tokios prieigos turi būti pridėtas atitinkamas sutikimas), arba
b) perkančioji organizacija ar perkantysis subjektas yra gavusi ir turi aktualius susijusius dokumentus iš ankstesnių (kitų) pirkimo procedūrų.

Ekonominės veiklos vykdytojai oficialiai sutinka perkančiajai organizacijai ar perkančiajam subjektui, nurodytam I dalyje, leisti susipažinti su dokumentais, kuriais patvirtinama informacija, pateikta šio Europos bendrojo viešųjų pirkimų dokumento III ir IV dalyse, kiek tai susiję su pirkimu, nurodytu I dalyje.

Data, vieta ir, jei reikia ar būtina, parašas (-ai):

Data

03-01-2025

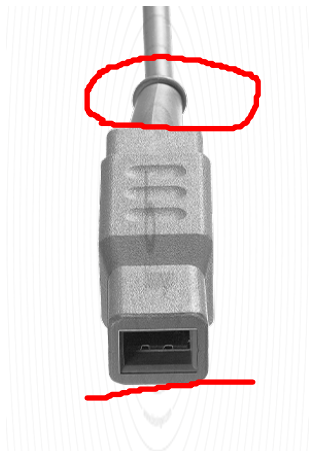
Vieta

Kaunas

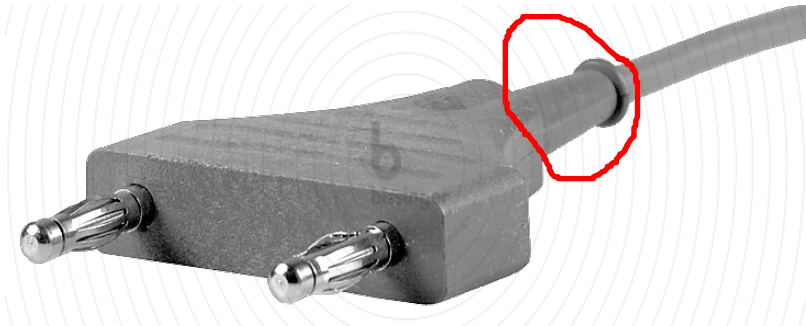
Parašas

80100033

Bipolar Cable



4.1.3 Su papildomomis apsaugomis nuo nulankstymo prie jungčių



4.1.5 Su europinio tipo plokščia jungtimi bipoliariams pincetams



Technical specifications

Dimensions and Weights

Cable length	5 m	4.1.4 Kabelio ilgis 5m
--------------	-----	------------------------

Basic Specifications

Cable connector instrument side	Flat Plug square
Cable plug generator side	Valleylab, Conmed, International

Cleaning and Reprocessing

Reusable	Yes	4.1.2 Daugkartinio naudojimo
Manual reprocessing	Yes, related to IFU	
Washing machine	95 °C	
Autoclave sterilization	max. 137 °C	

4.1.1 sterilizuojamas

Approvals

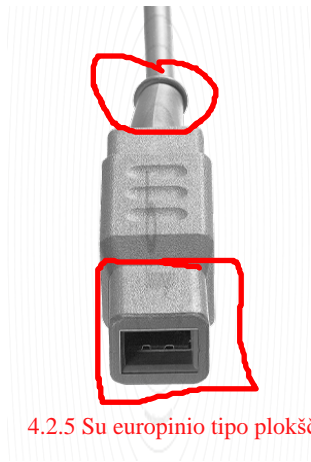
Directive 93/42 EEC	N/A
Regulation (EU) 2017/745	Class I
FDA Approval	Class II

Additional Product Information

Accessories rated voltage	max. 1 kVp
Packaging unit	1
GTIN 14	04250418700313

80100067

Bipolar Cable



4.2.3 Su papildomomis apsaugomis nuo nulankstymo prie jungčių



4.2.5 Su europinio tipo plokščia jungtimi bipolariniams pincetams



Technical specifications

Dimensions and Weights

Cable length	5 m	4.2.4 Kabelio ilgis 5m
--------------	-----	------------------------

Basic Specifications

Cable connector instrument side	Flat Plug square
Cable plug generator side	Erbe, EMC/Dolley

Cleaning and Reprocessing

Reusable	Yes	4.2.2 daugkartinio naudojimo
Manual reprocessing	Yes, related to IFU	
Washing machine	95 °C	
Autoclave sterilization	max. 137 °C	

4.2.1 Sterilizuojamas

Approvals

Directive 93/42 EEC	N/A
Regulation (EU) 2017/745	Class I
FDA Approval	Class II

Additional Product Information

Accessories rated voltage	max. 1 kVp
Packaging unit	1
GTIN 14	04250418732147

Naudojimo ir saugos nurodymai

Jei nepaisysite šių naudojimo ir saugos nurodymų, galite susižeisti, gali būti vykdomos netinkamos funkcijos ir įvykti netikėtų neigiamų įvykių.

- Visi kabeliai prieš naudojant ir po to turi būti visiškai nuvalyti, dezinfekuoti ir sterilizuoti.
- Kiekvieną kartą prieš naudojant kabelius reikia vizualiai patikrinti ir atlikti funkcijų testą.
- Įsitikinkite, kad pasirinktas kabelis yra su tinkamomis jungtimis tiek elektrochirurginiam generatoriui tiek naudojamam instrumentui. Taip pat patikrinkite ar abi jungtis pilnai įstatėte.
- Draudžiama naudoti pažeistus kabelius.
- Tam, kad išvengti sugadinimo, kabelio nelankstykite.
- Tam, kad išvengti kabelio pažeidimų, atjunginėkite kabelius juos prilaikydami prie elektrochirurginio generatoriaus kištumo.
- Nenaudokite, jei aplinkoje yra degių ar sprogių medžiagų.

Parngiamasis apdorojimas

Dėl gaminio dizaino, naudojamų medžiagų ir naudojimo paskirties negalima nustatyti tikslaus didžiausio galimų parngiamųjų apdorojimų ciklų limito. Instrumentų naudojimo trukmę lemia jų atliekamos funkcijos ir tinkamas naudojimas juos tausojant.

Paruošimas ir transportavimas

Kiekvieną kartą panaudojus, būtina nedelsiant nuo kabelio nuvalyti visus matomus nešvarumus. Nenaudokite fiksuojamųjų priemonių ar karšto vandens (> 40 °C).

Mašininis parngiamasis apdorojimas

Valymas

Instrumentus sudėkite ant įstumiamačio modulio sietinio padėklo arba ant mikrochirurginių prietaisų modulio ir pradėkite valymo procesą.

1. 1 min. skalaukite šaltu vandeniu.
2. Ištuštinkite.
3. 3 min. skalaukite šaltu vandeniu.
4. Ištuštinkite.
5. 5 min. plaukite 55 °C temperatūroje 0,5 % šarminiu valikliu arba 45 °C temperatūroje fermentiniu valikliu.
6. Ištuštinkite.
7. 3 min. neutralizuokite šiltu vandentiekio vandeniu (> 40 °C) ir neutralizatoriumi.
8. Ištuštinkite.
9. 2 min. įjunkite tarpinį skalavimą šiltu vandentiekio vandeniu (> 40 °C).
10. Ištuštinkite.

5.1.1, 5.1.2, 5.2.1, 5.2.2 p.
sterilizuojamas, daugkartinio naudojimo

Dezinfekcija

Mašininė šiluminė dezinfekcija, atsižvelgiant į šalyje galiojančius reikalavimus dėl A0 reikšmės (žr. ISO 15883).

Džiovinimas

Instrumentų išorės džiovinimas valymo / dezinfekavimo prietaise įjungus džiovinimo ciklą. Jei reikia, galima papildomai nusausinoti nepūkuota servetėle. Tuščias ertmes išsausinkite steriliu suslėgtuoju oru.

Rankinis parngiamasis apdorojimas

Valymas

Valymo vonelę paruoškite pagal gamintojo duomenis.

1. Gaminį skalaukite šaltu vandentiekio vandeniu (< 40 °C), kol pašalinsite visus matomus nešvarumus. Prilipusius nešvarumus pašalinkite minkštu šepetėliu.
2. Gaminį visiškai panardinkite į paruoštą valymo vonelę. Laikykites gamintojo nurodyto poveikio laiko.
3. Įdėtą instrumentą nuvalykite minkštu šepetėliu. Visus paviršius šepetėliu

nuvalykite kelis kartus.

4. Gaminį kruopščiai išskalaukite demineralizuotu vandeniu, kad pašalintumėte visus valymo priemonės likučius.

Dezinfekcija

Dezinfekavimo vonelę paruoškite pagal dezinfekavimo priemonės gamintojo duomenis. Sudėkite instrumentus į dezinfekavimo vonelę ir atkreipkite dėmesį į nurodytą poveikio laiką. Gaminį labai kruopščiai nuskalaukite demineralizuotu vandeniu, kad pašalintumėte visus dezinfekavimo priemonės likučius.

Džiovinimas

Rankomis nusausinama naudojant nepūkuotą servetėlę, o tuščios ertmės ir kanalai išsausinami steriliu suslėgtuoju oru.

Veikimo patikra ir pakuotė

Reikia apžiūrėti, ar instrumentai švarūs ir nepažeisti. Jei reikia, kartokite parengiamojo apdorojimo procesą, kol instrumentas taps švarus. Standartus atitinkanti sterilizuotų instrumentų pakuotė pagal ISO 11607 ir EN 868.

Sterilizavimas

Gaminiai sterilizuojami naudojant frakcinį pirminio vakuumavimo metodą (pagal ISO 13060 / ISO 17665), atsižvelgiant į atitinkamus šalyje galiojančius reikalavimus.

- Per 3 pirminio vakuumavimo fazes naudojamas mažiausiai 60 mbar slėgis.
- Įkaitinama mažiausiai iki 132 °C sterilizavimo temperatūros; didžiausia temperatūra – 137 °C.
- Laikymo laikas: mažiausiai 3 min.; ilgiausiai 18 min.
- Džiovinimo laikas: mažiausiai 10 min.

⚠ Jeigu yra įtarimas dėl prionų infekcijos (CLI) tuomet, vadovaujantis nacionalinėmis gairėmis, galimas ilgesnis laikymo laikas (pvz., 15 min.).

Remontas

Neremontuokite patys. Techninės priežiūros ir remonto darbus gali atlikti tik atitinkamai išmokyti ir kvalifikuoti asmenys. Jei kiltų susijusių klausimų, kreipkitės į gamintoją arba savo medicinos technikos skyrių.

⚠ Prieš siunčiant remontuoti sugedusius gaminius, reikia juos išvalyti, taikant parengiamojo apdorojimo procedūras.

Informacija dėl parngiamojo apdorojimo kontrolės

Atliekant kontrolę buvo naudojamos tokios patikros instrukcijos, medžiagos ir mašinos:

Valymo priemonės (mašininiam valymui):

Neodisher FA; „Dr. Weigert“ (šarminis)

Endozime, bendrovė „Ruhof“ (fermentinis)

Valymo priemonės (rankiniam valymui):

„Cidezyme“, „Enzol“ fermentinis detergantas, „Johnson & Johnson“

Dezinfekavimo priemonės (rankiniam valymui):

„Cidex OPA“, „Johnson & Johnson“

Neutralizatorius:

„Neodisher Z“; „Dr. Weigert“

Valymo ir dezinfekavimo prietaisai:

„Miele“ dezinfekatorius „G 7735 CD“

„Miele“ įstumiamačio modulis „E 327-06“

„Miele“ mikrochirurginių prietaisų modulis „E 450“

Smulkesnė informacija prieinama pranešime.

SMP GmbH # 01707011901-2

MDS GmbH # 135196-10

Nelson Labs # 200432706-02

MDS GmbH Testbericht 084183-10

(automatinis valymas)
(rankinis valymas, sterilizacija)
(sterilizacija)
(sterilizacija)



Stecker generatorseitig
Plug, generator side
Connecteur, côté générateur
Enchufe, lado del generador
Spina, lato generatore

Flachstecker Flat plug Connecteur plat Enchufe plano Spina piatta	Flachstecker gewinkelt Flat plug, angled Connecteur plat, coudé Enchufe plano acodado Spina piatta, ad angolo	Flachstecker Vierkant Flat plug, square Connecteur plat, carré Enchufe plano cuadrado Spina piatta, quadrata	2-Pin-Stecker 2-pin plug Connecteur à 2 broches Enchufe 2 pines Spina a 2 pin	2-Pin-Stecker gewinkelt 2-pin plug, angled Connecteur à 2 broches, coudé Enchufe 2 pines acodado Spina a 2 pin, ad angolo	2-Pin-Stecker gewinkelt mit Schutzkappe 2-pin plug, angled with protecting cap Connecteur à 2 broches, coudé avec capuchon de protection Enchufe 2 pines acodado con caperuza protectora Spina a 2 pin, ad angolo con cappuccio di protezione	Rundstecker BiTech, WAVE, Plasmaloop Round plug BiTech, WAVE, Plasmaloop Connecteur rond BiTech, WAVE, Plasmaloop Enchufe redondo BiTech, WAVE, Plasmaloop Spina rotonda, BiTech, WAVE, Plasmaloop	Stecker Zange R. Wolf Plug R. Wolf forceps Connecteur, pince R. Wolf Enchufe pinza R. Wolf Spina, pinza R. Wolf	Stecker Zange Karl Storz Plug, Karl Storz forceps Connecteur, pince Karl Storz Enchufe pinza Karl Storz Spina, pinza Karl Storz	Stecker Ethicon Scheren, Scheren-Klemmen Plug Ethicon scissors, clamp-scissors Connecteur, ciseaux, ciseaux à griffes Ethicon Enchufe tijeras, pinzas- tijeras Ethicon Spina, forbici, forbici a pinza Ethicon	Stecker Aesculap Pinzetten Plug, Aesculap forceps Connecteur, pincas Aesculap Enchufe pinzas Aesculap Spina, pinze Aesculap	Stecker Storz Resektoskop Plug, Storz resectoscope Connecteur, Résectoscope Storz Enchufe Resectoscopio Storz Spina, Resettoscopio Storz	Stecker Wolf Resektoskop Plug, Wolf resectoscope Connecteur, Résectoscope Wolf Enchufe Resectoscopio Wolf Spina, Resettoscopio Wolf

*Entspricht nicht
EU-Bestimmungen

*Non-compliant with
EU standards

*Non conforme aux
normes UE

*No cumple las
normas UE

*Non conforme alle
norme UE

 2-Banana*	3 m	80100010	80100051	80100012	80100025	80100125	80100128	80100081	80100094	80100098	80100200	80100121	-	-
	5 m	80100013	80100052	80100011	80100027	80100127	80100129	80100085	-	80100201	80100300	80100122	-	-
 U20	4 m	80100260	80100261	80100262	80100263	80100264	80100265	80100266	80100267	80100268	80100269	80100270	-	-
 Erbe	3 m	80100017	80100054	80100037	80100023	-	-	80100082	80100092	80100095	80100198	80100150	-	-
	5 m	80100018	80100056	80100038	80100029	80100132	-	80100086	80100192	80100195	80100298	80100169	-	-
 Martin, Berchtold, Aesculap GK 55, GK 60	3 m	80100019	80100058	80100039	80100043	-	-	80100083	80100093	80100096	80100199	80100130	-	-
	5 m	80100020	80100059	80100047	80100045	-	-	80100087	80100193	80100196	80100299	80100131	-	-
 Valleylab, Lamidey, EMC	3 m	80100021	80100046	80100034	80100053	80100153	80100152	80100084	80100089	80100097	80100197	80100170	-	-
	4 m	-	-	-	-	-	-	-	-	-	-	-	80100276	80100277
	5 m	80100022	80100048	80100033	80100055	80100155	80100154	80100088	80100189	-	80100297	80100171	-	-
 Erbe, EMC/Dolley	3 m	80100050	80100061	80100057	80100044	80100064	80100156	80100080	-	80100099	-	-	-	-
	5 m	80100060	-	80100067	-	-	-	80100104	-	-	-	-	-	-
 Valleylab autobipolar	3 m	80100042	-	80100219	-	-	-	-	-	-	-	-	-	-
	5 m	-	-	-	-	-	-	-	-	-	-	-	-	-
 Ellman	4 m	-	-	80100274	80100275	-	-	-	-	-	-	-	-	-

4.1.6.Su 29 mm pločio 2
kontaktų tarptautinė
jungtimi pajungimui prie
generatoriaus

4.2.6 .Su 22mm pločio 2
kontaktų tarptautinė
jungtimi pajungimui prie
generatoriaus



Stecker generatorseitig
Plug, generator side
Connecteur, côté générateur
Enchufe, lado del generador
Spina, lato generatore



Ø 4 mm
Buchse
Socket
Douille
Zócalo
Bussola



Ø 4 mm
Buchse
gewinkelt
Socket, angled
Douille, coudée
Zócalo acodado
Bussola ad angolo



Ø 2 mm
Stecker geschützt
für Resektoskop
**Plug, protected
for resectoscope**
Connecteur protégé
pour résectoscope
**Enchufe
protegido para
el resectoscopio**
*Spina protetta
per resettoscopio*



Ø 4 mm
Stecker
geschützt
Plug, protected
Connecteur
protégé
**Enchufe
protegido**
Spina protetta



Ø 3 mm
Buchse
Socket
Douille
Zócalo
Bussola



Ø 4 mm
Buchse
mit Sechskant
Hexagon socket
Douille hexagonale
**Zócalo
con hexágono**
*Bussola
con esagono*



Ø 4 mm
Stecker
Plug
Connecteur
Enchufe
Spina



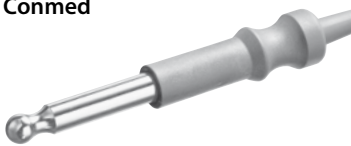
Ø 4 mm
Buchse mit
Schutzhülse
**Socket with protec-
tive sleeve**
Douille avec man-
chon de protection
**Zócalo con la
manga protectora**
*Bussola con mani-
cotto protettivo*

Erbe T-Serie, Martin,
Berchtold



3 m	80100014	80100240	80100314		-	80100319	80100315	80100214	-
4 m	-	-	-		-	-	-	-	80100015
5 m	80100016	80100241	80100324		-	-	-	80100216	-

Bovie, Valleylab,
Conmed



3 m	80100215	80100242	80100320		80100220	80100322	80100312	80100217	-
4 m	-	-	-		-	-	-	-	80100251
5 m	80100221	80100245	80100340		80100222	-	-	80100218	-

Martin, Berchtold








3 m	80100031	-	80100316		80100230	-	-	80100212	-
5 m	80100331	-	80100326		-	-	-	-	-

Erbe ACC/ICC



3 m	80100114	80100243	80100330		-	-	80100318	-	-
4 m	-	-	-		-	-	-	-	80100252
5 m	80100115	-	80100360		-	80100317	-	-	-

Kabel für Einmal-Neutralelektroden
Cables for disposable neutral electrodes
Câbles pour électrodes neutres à usage unique
Cable para electrodos neutros desechables
Cavi per elettrodi neutri monouso

Stecker generatorseitig Plug, generator side Connecteur, côté générateur Enchufe, lado del generador Spina, lato generatore		 Stecker für Einmal-Neutral-Elektroden Plug for single-use electrodes Connecteur pour électrodes neutres à usage unique Enchufe para electrodo desechable Spina per elettrodi neutri monouso	
	4,5 m	89101050	
	4,5 m	89101052	
	4,5 m	89101051	
	4,5 m	89101053	

5.1.5 Prijungimas – 2 kontaktų REM tarptautinis

5.2.5 Prijungimas – 2 kontaktų NON REM tarptautinis

5.1.3 ir 5.2.3 Su papildomomis apsaugomis nuo nulankstymo prie jungčių

5.1.4 Kabelio ilgis 4,5 m;

5.2.4 kabelio ilgis 4,5m

Die wiederverwendbaren Neutral-elektroden von Bissinger bestehen aus leitfähigem, widerstandsfähigem Silikon und sind mit einem 4 m Anschlußkabel ausgestattet.

The reusable neutral electrodes by Bissinger are made of conductive resistant silicone and are equipped with a 4 m connecting cable.

Les électrodes neutres réutilisables de Bissinger sont faites de silicone conductrice résistante et dotées d'un câble de raccordement de 4 m.

Los electrodos neutros reutilizables de Bissinger se componen de silicona conductiva y resistente y están equipados con un cable de conexión de 4 m.

Gli elettrodi neutri riutilizzabili della Bissinger sono fatti da silicone conduttivo e resistente e sono muniti di un cavo di collegamento di 4 m.

Wisch- oder Sprühdesinfektion
Wipe or spray disinfection
Désinfection par essuyage ou pulvérisation
Desinfección por lavado o rocío
Disinfezione mediante pulizia o con spray

*Entspricht nicht EU-Bestimmungen
*Non-compliant with EU standards
*Non conforme aux normes UE
*No cumple las normas UE
*Non conforme alle norme UE

Wiederverwendbare Neutralelektroden
Reusable neutral electrodes
Électrodes neutres réutilisables
Electrodos neutros reutilizables
Elettrodi neutri riutilizzabili



Kontaktfläche 390 cm² (15 x 26 cm), für Erwachsene
Contact surface 390 cm² (15 x 26 cm), for adult
Surface de contact 390 cm² (15 x 26 cm), pour adultes
Superficie de contacto 390 cm² (15 x 26 cm), para adultos
Superficie di contatto 390 cm² (15x26 cm), per adulti

Stecker generatorseitig
Plug, generator side
Connecteur, côté générateur
Enchufe, lado del generador
Spina, lato generatore

	89242004*
	89242003*
	89242005*

Kontaktfläche 128 cm² (8 x 16 cm), für Kinder
Contact surface 128 cm² (8 x 16 cm), for children
Surface de contact 128 cm² (8 x 16 cm), pour enfants
Superficie de contacto 128 cm² (8 x 16 cm), para niños
Superficie di contatto 390 cm² (15x26 cm), per bambini

Stecker generatorseitig
Plug, generator side
Connecteur, côté générateur
Enchufe, lado del generador
Spina, lato generatore

	89232004*
	89232003*
	89232005*

VALSTYBĖS ĮMONĖ REGISTRŲ CENTRAS

Studentų g. 39, 08106 Vilnius, tel. +370 5 268 8262, el. p. info@registrucentras.lt

**KOMPETENTINGŲ INSTITUCIJŲ TVARKOMŲ JUNGTINIŲ DUOMENŲ APIE VIEŠŲJŲ
PIRKIMŲ PROCEDŪROJE DALYVAUJANTĮ TIEKĖJĄ (JURIDINĮ ASMENĮ)
PAŽYMA**

2025-03-17 Nr. 762235

Tiekėjo pavadinimas	UAB "Sorimpeksas"
Tiekėjo kontaktinė informacija:	
mobilusis telefonas	+37069825117
elektroninio pašto adresas	info@sorimpeksas.com
interneto svetainės adresas	www.sorimpeksas.com
Buhalterio (buhalterių) ar kito (kitų) asmens (asmenų), turinčio (turinčių) teisę surašyti ir pasirašyti tiekėjo apskaitos dokumentus, vardas, pavardė	
<u>Juridinių asmenų registras:</u>	
kodas	135733248
teisinė forma	Uždaroji akcinė bendrovė
teisinis statusas	Teisinis statusas neįregistruotas
buveinė (adresas)	Kaunas, Šiaulių g. 16A, LT-44353
Vadovo, kito valdymo ar priežiūros organo nario ar kito asmens, turinčio (turinčių) teisę atstovauti tiekėjui ar jį kontroliuoti, jo vardu priimti sprendimą, sudaryti sandorį, vardas, pavardė	RAMŪNAS ŽALNERAUSKAS
įregistravimo data	2001-03-02
<u>Valstybinė mokesčių inspekcija prie Lietuvos Respublikos finansų ministerijos:</u>	
duomenys apie tiekėjo atsiskaitymą su valstybės, savivaldybių biudžetais ir valstybės pinigų fondais	Atsiskaitęs
Duomenų suformavimo data	2025-03-13
<u>Valstybinio socialinio draudimo fondo valdyba prie Socialinės apsaugos ir darbo ministerijos:</u>	
duomenys apie tiekėjo atsiskaitymą su Valstybinio socialinio draudimo fondu	Neįsiskolinęs
Duomenų suformavimo data	2025-03-13
<u>Įtariamųjų, kaltinamųjų ir nuteistųjų registras:</u>	
duomenys apie tiekėją	Dėl UAB "Sorimpeksas", kodas 135733248, per pastaruosius 5 metus nėra priimtas ir įsiteisėjęs apkaltinamasis teismo nuosprendis už nusikalstamas veikas, nurodytas Lietuvos Respublikos viešųjų pirkimų įstatymo 46 straipsnio 1 dalyje ir 3 dalyje. Nėra paskirta baudžiamojo poveikio priemonė - uždraudimas juridiniam asmeniui dalyvauti viešuosiuose pirkimuose pagal Viešųjų pirkimų įstatymo 46 straipsnio 2-1 dalį.

duomenys apie tiekėjo vadovą, kitą valdymo ar
priežiūros organo narį ar kitą (kitus) asmenį
(asmenis), turintį (turinčius) teisę atstovauti
tiekėjui ar jį kontroliuoti, jo vardu priimti
sprendimą, sudaryti sandorį

duomenys apie tiekėjo buhalterį (buhalterius) ar
kitą (kitus) asmenį (asmenis), turintį (turinčius)
teisę surašyti ir pasirašyti tiekėjo apskaitos
dokumentus

Duomenų suformavimo data

**Ramūnui Žalnerauskui, gim. 1970 m. lapkričio 13 d., per
pastaruosius 5 metus nėra priimtas ir įsiteisėjęs
apkaltinamasis teismo nuosprendis ir jis neturi
neišnykusio ar nepanaikinto teistumo už nusikalstamas
veikas, nurodytas Lietuvos Respublikos viešųjų pirkimų
įstatymo 46 straipsnio 1 dalyje.**

**[redacted] per
pastaruosius 5 metus nėra priimtas ir įsiteisėjęs
apkaltinamasis teismo nuosprendis ir ji neturi neišnykusio
ar nepanaikinto teistumo už nusikalstamas veikas,
nurodytas Lietuvos Respublikos viešųjų pirkimų įstatymo
46 straipsnio 1 dalyje.**

2025-03-14

Pažymą išspausdino:

Asmenų registravimo centro Juridinių asmenų registro
Kauno skyriaus Kauno 1 Juridinių asmenų registro
duomenų tvarkymo grupės
Registratorė

[redacted]

A. V.



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
ZLG-BS-244.10.08



Product Service

EC Certificate

Full Quality Assurance System

Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)
(Devices in Class IIa, IIb or III)

No. G1 077608 0079 Rev. 00

Manufacturer:

Covidien llc

15 Hampshire Street
Mansfield, MA 02048
USA

Product Category(ies): Medical Instruments, Surgical Products
and Hemostatic Materials:

- Surgical Suture Products, Pledgets and Retention Tapes
- Endoscopy Instruments and
Accessories including Lubricant
- Surgical Staple, Clip Products and Accessories
- Manual Surgical Instruments
- Implantable Wound Dressing Materials
- Ultrasonic Surgical Devices and Accessories
- Suction / Irrigation Devices and Accessories
- Arthroscopy Implants, Instruments and Accessories
- Bone Wax
- Temporary Cardiac Pacing Lead
- Powered Stapling Systems

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device categories in accordance with MDD Annex II. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of class III devices an additional Annex II (4) certificate is mandatory. See also notes overleaf.

Report No.:

713164286

Valid from:

2019-09-13

Valid until:

2024-05-26

Date,

2019-09-13



Page 1 of 2

TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany

TÜV®

ZERTIFIKAT • CERTIFICATE • 認證證書 • CERTIFICADO • CERTIFICAT

EC Certificate

Full Quality Assurance System

**Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)
(Devices in Class IIa, IIb or III)**

No. G1 077608 0079 Rev. 00

Facility(ies):

Covidien Inc
15 Hampshire Street,
Mansfield, MA 02048, USA

4

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

Manufacturer's Declaration

in relation to Regulation (EU) 2023/607 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices, in particular with respect to

- the validity of certificates issued under Council Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD) or Council Directive 93/42/EEC on Medical Devices (MDD) (Directive Certificates) *and/or*¹
- the compliance of the devices and us as their manufacturer with the conditions for the continued placing on the market and putting into service

Manufacturer name	Covidien llc
Manufacturer address and contact details	15 Hampshire Street Mansfield, MA 02048 USA
Single Registration Number (SRN) (if available)	US-MF-000028763

Authorised Representative name (if applicable)	Covidien Ireland Limited
Authorised Representative address and contact details	IDA Business and Technology Park Tullamore, Ireland
Single Registration Number (SRN) (if available)	

Notified body name (if applicable)	<input checked="" type="checkbox"/> See attached schedule
Notified body number (if applicable)	<input checked="" type="checkbox"/> See attached schedule
Directive Certificate number(s) to which this confirmation is made (if applicable)	<input checked="" type="checkbox"/> See attached schedule
Original expiry date as indicated on the Directive Certificate prior to the extension of the validity (if applicable)	<input checked="" type="checkbox"/> See attached schedule
End date of extended validity/transition period	<input checked="" type="checkbox"/> See attached schedule

¹ The first condition is not applicable in case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body.

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

We, as the manufacturer declare under our sole responsibility:

- for the above listed **Directive Certificate** (or see attached schedule, if multiple certificates) the conditions for the legal extension of validity as required in Article 120.2 of the MDR are met *and/or*²
- the listed **device(s)** in the attached schedule and we as their manufacturer are in compliance with the conditions listed in Article 120.3c of the MDR for continued placing on the market and putting into service,

namely by fulfilling the following conditions:

➤ **Directive Certificate(s)** as listed above or in the attached schedule

- Directive Certificate(s) covering the listed device(s) was/were issued after 25 May 2017, was/were valid on 26 May 2021 and have not been withdrawn afterwards.

Choose applicable statements:

☐ Expired *before* 20 March 2023:

- ☐ Before the original date of expiry as indicated on the Directive Certificate(s), we and the notified body have signed written agreement(s) in accordance with Section 4.3, second subparagraph of Annex VII to this Regulation for the conformity assessment(s) in respect of the device(s) covered by the expired certificate(s) or in respect of a device(s) intended to substitute that/those device(s), or
- ☐ A Competent Authority has granted a derogation from the applicable conformity assessment procedure in accordance with Article 59(1) MDR (may be provided upon request), or
- ☐ A Competent Authority has required the manufacturer, in accordance with Article 97(1) MDR, to carry out the applicable conformity assessment procedure (may be provided upon request)

Choose one of the following statements only if a derogation per Article 59(1) or a requirement per Article 97(1) has been granted by a Competent Authority:

- ☐ Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitute(s) and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.
- ☐ We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

² The first condition is not applicable in case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

- ☒ Expired/expires *after* 20 March 2023:

Choose one applicable statement:

☒ Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitute(s) and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.

- ☐ We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

➤ **Upclassified devices**

In case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body:

Choose one applicable statement:

- ☐ Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitutes and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.
- ☐ We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

➤ **Quality Management System (QMS)**

Choose one applicable statement:

- ☐ A QMS in accordance with Article 10(9) MDR will be put in place by no later than 26 May 2024.
- ☐ A QMS in accordance with Article 10(9) MDR is in place.
- ☒ A notified body has issued the attached certificate for the MDR-compliant QMS.

➤ **Device(s) as listed in the attached schedule**

- The device(s) continue to comply with the AIMDD or MDD.
- There are no significant changes in the design and intended purpose.
- The device(s) do not present an unacceptable risk to health or safety of patients, users or other persons, or to other aspects of the protection of public health.

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

Signed for and on behalf of the manufacturer:

Full Company Name Covidien llc
Location 15 Hampshire Street, Mansfield, MA 02048 USA
Date: 22 February 2024

[Redacted signature]

[Redacted text]

[Redacted text]
Contact Details (at least email) [Redacted text]

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

Signed for and on behalf of the manufacturer:

Full Company Name Covidien Inc

Location 15 Hampshire Street, Mansfield, MA 02048 USA

Date: 22 February 2024

[REDACTED]

[REDACTED]

[REDACTED]
[REDACTED]
[REDACTED]

Contact Details (at least email) [REDACTED]

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

Schedule of Devices

The above Manufacturer's Declaration is valid for the following devices:

Identification of the device(s) ³ (e.g., device name, family/group name, device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)
133650	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
134048	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
134051	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
134053	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
150462	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
170001	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170002	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170003	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170004	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170050	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170051	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170052	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170053	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170054	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170055	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170056	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

170057	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170070	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170071	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170072	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170073	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170090	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170092	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170094	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
170096	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
173016	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173019	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173021	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
173022	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173023	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
173024	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
173026	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
173030	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173046	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173049	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173052	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
173054	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
174001	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
174006	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
174007	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
174015	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
174209	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
174301	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
174317	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

174601	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176605	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176613	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176625	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176630	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
176643	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176645	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176647	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176657	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179301	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179303	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179305	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179307	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179308	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179309	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
179310	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
8886262741	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886306261	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886337041	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886400241	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886400963	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886402721	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886402821	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886402831	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886403231	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440013	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440023	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440113	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886440123	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440133	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440223	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440233	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440243	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440333	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440343	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886440861	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441003	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441013	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441023	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441431	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441543	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441853	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441913	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441923	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886441933	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442013	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442023	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442033	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442043	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442153	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442233	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442243	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442253	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442431	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442441	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442451	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886442461	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442531	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442641	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442651	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886442661	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886443371	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886443671	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445041	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445241	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445251	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445261	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445463	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445473	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445561	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445571	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445863	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445951	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445961	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886445971	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886446063	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886600111	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886600741	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886600751	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886601563	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886603433	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886603443	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886605941	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886606163	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886606173	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886606773	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886611511	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886611831	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886612221	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886613111	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886613211	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614331	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614721	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886620133	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621311	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621321	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621331	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886622933	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886622943	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886622953	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623131	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623241	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623331	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623341	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623351	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623531	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623541	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623741	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886623761	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886624541	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886624551	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886624561	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886624571	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886624753	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886624961	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886625741	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886625751	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886625761	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626131	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626141	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626151	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626161	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626741	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626751	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626761	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626771	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626851	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626951	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626961	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886626971	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886627553	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886627563	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886627573	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886627761	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886628561	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886628571	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886628751	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886628761	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886628771	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886629571	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886630761	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886630771	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886630972	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886631163	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886631173	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886631962	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886631972	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886632271	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886633961	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886633971	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886639053	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886640351	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886640451	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886641971	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886645561	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886645661	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886653211	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886653221	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886653311	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886660521	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886660531	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886660821	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886660831	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886660841	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886661231	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886661241	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886661831	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886661841	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886661851	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886662451	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886662651	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886662801	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886663241	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886663931	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886663941	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886664141	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886664151	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886674441	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886674451	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886674551	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886675551	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886675561	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886675571	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886675771	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886676971	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886678771	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886690831	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886690841	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886803512	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
8886803712	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
8886848700	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
8886848701	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
8886848808	G70776080083	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886848812	G70776080083	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886848813	G70776080083	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886848882	G70776080083	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

020250	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
020251	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
173050G	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
176630B	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
21-345	G2S0776080072	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
220-50	G2S0776080072	5/23/2024	TUV 0123	TUV 0123	12/31/2028	
250-30	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
250-40	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
2-CL-852	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
31-121	G2S0776080072	5/23/2024	TUV 0123	TUV 0123	12/31/2028	
3-CL-812	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
3-CP-360	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861720-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861727-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861727-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861741-11	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861741-21	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861741-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861742-21	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861742-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861742-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861744-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861744-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861744-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861747-11	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861749-24	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861750-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861754-13	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88861756-11	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861756-21	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861756-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861756-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861756-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861757-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861757-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861764-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861764-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861799-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861799-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861799-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861883-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861883-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861883-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861895-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861896-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861899-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861899-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861902-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861902-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861906-42	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861906-52	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861909-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861910-22	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861910-32	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861913-72	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861914-83	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88861915-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861915-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861917-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861917-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861917-82	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861918-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861918-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861918-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861919-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861919-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861919-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861919-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861919-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861920-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861920-72	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861920-93	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861922-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861926-42	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861927-32	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861927-42	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861928-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861928-72	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861931-43	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861932-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861934-53	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861935-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861935-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861935-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88861948-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861948-81	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861951-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861951-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861951-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861953-32J	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861954-32J	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861957-32	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861958-42	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861958-52	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861962-52	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861962-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861962-72	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861964-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861964-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861967-52	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861968-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861969-52	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861969-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861969-72	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861971-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861971-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861981-33	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861982-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861985-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861985-61	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861985-71	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861989-32	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88861989-42	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861990-42J	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861991-42	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861993-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861995-31	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861995-41	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861998-62	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88861998-72	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862011-51	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862222-69	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862222-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862224-09	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862224-49	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862224-69	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862224-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862226-69	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862226-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862228-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862345-53	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862348-53	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862376-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862382-83	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862392-49	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862392-69	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862393-69	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862393-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862396-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862407-43	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88862407-49	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862407-53	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862407-63	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862408-73	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862408-83	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862410-69	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862410-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862412-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862413-83	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862414-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862420-09	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862420-83	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862424-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862466-25	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862494-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862495-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862497-89	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862631-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886263656-2	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886265856	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862768-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862775-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862775-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862775-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862801-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862808-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862811-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862815-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88862818-89	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862839-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862840-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862851-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862852-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862854-33	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862855-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862862-34	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862877-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862879-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862880-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862880-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862914-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862914-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862947-53	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862953-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862962-83	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862978-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88862981-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863001-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863001-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863001-72	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863002-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886300-32	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863003-32	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863003-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863008-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863008-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863008-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863008-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863011-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863012-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863015-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863017-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863017-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863023-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886302361	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863023-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863026-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863026-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863027-79	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863028-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863028-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863033-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863035-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863035-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863037-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863040-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863045-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863046-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863047-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863047-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863048-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863050-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863050-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863050-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863052-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863054-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863054-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863054-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863054-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863055-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863056-89	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863059-53	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863062-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863062-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863065-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863065-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863069-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863070-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863082-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863082-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863083-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863084-46	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863084-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863086-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863087-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863087-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863087-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863088-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863088-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863090-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863090-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863090-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863090-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863091-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863092-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863092-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863093-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863093-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863097-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863097-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863103-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886310421-2	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863106-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863107-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863109-43	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863111-79	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863113-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863113-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863119-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863125-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863127-79	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863128-79	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863128-82	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863129-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863139-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863146-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863147-83	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863147-89	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863154-09	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863155-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863157-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863159-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863159-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863159-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863160-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863160-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863160-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863163-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863163-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863163-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863165-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863167-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863167-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863167-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863167-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863168-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863182-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863182-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863182-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863185-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863185-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863185-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863185-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863185-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863186-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863186-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863186-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863186-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863190-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863191-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863193-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863202-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863202-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863203-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863204-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863205-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863205-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863205-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863211-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863212-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863213-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863213-66	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863218-36	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863218-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863219-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863220-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863221-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863222-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863226-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863226-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863226-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863227-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863227-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863228-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863229-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863229-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863230-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863233-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863235-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863236-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863239-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863240-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863241-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863249-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863251-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863255-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863256-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863256-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863256-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863256-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863257-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863258-32	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863258-42	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863258-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863259-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863259-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863260-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863260-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863260-72	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863261-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863261-72	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863261-82	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886326742-2	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863269-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863271-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863271-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863271-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863273-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863274-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863274-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863275-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863277-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863280-11	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863280-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863280-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863280-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863290-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863295-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863297-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863298-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863303-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863306-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863307-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863309-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863309-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863309-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863314-21	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863314-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863319-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863323-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863324-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863324-66	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863336-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863336-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863344-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863348-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863348-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863353-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863356-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863365-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863369-31	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863369-41	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863369-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863370-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863372-72	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863372-82	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863374-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863374-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863375-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863376-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863379-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863379-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863380-42	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863380-52	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863380-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863380-72	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863380-82	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863381-09	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863381-89	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863385-56	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88863386-62	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863393-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863393-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863395-51	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863395-61	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863395-71	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88863395-81	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864458-53	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864463-71	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864464-73	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864465-71	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864465-81	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864469-61	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864469-71	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864470-71	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864470-81	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864490-41	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864490-51	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864538-63	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864554-41	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864555-23	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864582-43	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864582-53	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88864619-51	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
8886603641-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886605731-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886612111-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886612411-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

8886612421-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886612701-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886612711-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886612821-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886613431-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886613901-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886613911-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614501-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614711-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614721-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614811-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614821-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886614921-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886616523-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886620721-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886620731-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886620741-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621721-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621731-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621741-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621933-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886621943-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886622131-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866277-71	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866288-11	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866288-21	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866292-51	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866297-11	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

88866297-21	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886646221-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
8886646231-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866751-71	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88866768-11	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
88868675-01	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88868677-01	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88868678-01	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88868679-01	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
88868682-01	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ABSTACK15	G70776080073	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ABSTACK20S	G70776080073	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ABSTACK30	G70776080073	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ABSTACK30X	G70776080073	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
B11STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B11STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B12LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B12LGS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B12STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B12STFCS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B5SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B5SHS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B5STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
B5STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
BPT12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
BZ4212	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
C-212	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

C-214	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-1945G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-196	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-543	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-658	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-801	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-802	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-803	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-810	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-811	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC811G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-812	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-813	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC813G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-830	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-831	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-843	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-844	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-845	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-872	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-883	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-884	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-885	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-904	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-905	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC905G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-912	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-913	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CC913G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-914	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-915	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-922	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-923	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-924	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-925	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-975	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CC-983	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CD-0423K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CD-3113K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CD-3123K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-005	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-015	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-030	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-045	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-047	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-055	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-060	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-10-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-11-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-12-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-13-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-14-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-15-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-16-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-17-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-200-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-201-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-202-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-20-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-219	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-21-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-22-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-232	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-239	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-240-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-250-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-259	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-270	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-271	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-2M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-304	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-30-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-31-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-330	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-331	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-354-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-355-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-356-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-358-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-35-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-369-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-36-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-370-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-374-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-375-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-376-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-377-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-380-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-381-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-382-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-383-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-42-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-457	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-460	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-461	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-462	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-4M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-510	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-511	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-513	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-515	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-516	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-517	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-526	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-528	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-52-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-53	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-533	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-534	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-535	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-536	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-537	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-538	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-53-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-542	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-543	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-544	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-545	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-547	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-552	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-554	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-555	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-55-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-568	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-569	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-570	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-576	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-581	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-595	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-596	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-597	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-5-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-602	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-603	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-60-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-622-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-623-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-63	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-633-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-644-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-64-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-659-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-665-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-673	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-673G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-677	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL6G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-6-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-71-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-730	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-739	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-740-L	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-741-L	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-769	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-779	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-782	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-800	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-801	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-802	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-803	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-804	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-805	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-809	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-810	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-811	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-812	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-813	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-814	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-815	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-816	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-817	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-818	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-823	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-824	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL824L	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-825	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL829	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-82-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-830	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-831	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-832	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-833	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-835-L	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-837	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-838	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-839	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-84	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-840	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-841	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-842	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-843	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-844	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-845	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-85	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-851-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-853	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-855	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-862	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-863	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-864	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-865	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-866	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-867	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-868	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-869	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-86-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-870	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-871	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-871-L	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-87-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-880	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-882	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-883	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-884	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-885	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-885-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-886-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-892	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-893	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-894	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-8-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-903	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-904	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-905	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-906	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-911	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-912	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-913	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-914	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-915	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-916	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-917	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-918	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-919	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-922	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-923	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-924	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-925	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-926	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-927	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-928	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-929	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-930	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-931	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-932	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-933	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-934	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-936	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-937	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-939	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-940	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-941	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-942	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-943	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-944	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-945	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-946	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-947	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-948	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-949	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-950	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-951	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-952	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-953	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-954	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-955	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-956	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-957	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-958	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-962	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-963	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-964	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-965	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-968	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-969	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-970	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-971	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-972	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-973	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-974	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CL-975	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-976	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-977	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-979	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-97-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-982	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-990	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-992	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-996-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-997-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-998-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CL-9-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-11-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-13-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT151MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-15-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-20-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-21-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-250-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-259	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-623-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-633-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-844	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-850-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-851-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-855	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-952	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-953	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CLT-954	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-955	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-957	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-958	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CLT-9-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CMM1972G	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
CMM-435-MG	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
CN-225	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-490	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-623	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-722	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-723	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN723G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-724	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-771	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-791	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-792	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-793	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN793G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-819	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-822	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-823-L	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-824	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-825	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-825-L	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN830GL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-830-L	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CN-925	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CN-926	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-10-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-16-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-17-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-30-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-411	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-412	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-414	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-415	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP41G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-422	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-423	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-424	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-425	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-433	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-434	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-435	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-443	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-453	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-454	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-455	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-460	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-535	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-824	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-824-L	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-825	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-834-L	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-836	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CP-844	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-845	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-92-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CP-957	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CPB-738G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-10-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-16-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-17-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-210	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-211	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-390	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-423	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-424	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-425	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-482	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-490	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-562	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-744	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-745	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-746	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-791	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-792	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS792G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-793	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS793G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-794	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-85-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
CS-92-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

CS-9-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1683K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1722K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1760K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1763K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1764K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1779K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1780K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
D-1879K	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
E0510	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E0512	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1020	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450-4	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450-4NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450-6	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450-6NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450G	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450G-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450X	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1450XNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1452	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1452-6	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1452NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455-4	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455-4NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455-6	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455-6NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

E1455B	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455B-4	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455B-4NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455BNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455G	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1455NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465-4	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465-4NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465-6	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465B	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465B-4	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465-BNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1465NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1475X	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1502	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1504	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1550	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1551-6	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1551-6-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1551G	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1551X	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1551X-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1552	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1552-6	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1552-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1559	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1560	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

E1561	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1562	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1563	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1564	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1565	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1567	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1650	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1651	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1652	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1653	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E1654B	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2100	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2100E	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2350H	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2350H-DB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2350H-DBNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2350HNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2400	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2401	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2450H	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2450HDBNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2450H-GNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2450HNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2505-10FR	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2505-10FRNSB	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515GNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515H	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

E2515H-DA	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515HDANSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515HDB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515H-DB-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515H-GNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515H-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2515-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516-GNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516H	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516H-DA	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516H-GNSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516H-NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2516NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2608-6	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2750	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E2750NSB	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3250H	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3250HNSB	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3305	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3310	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3310NSB	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3504	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3504H	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3504NSB	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3770-36	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3770-36C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3771-36	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

E3771-36C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3771-45	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3771-45C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3772-36	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3772-36C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3773-36	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3773-36C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3773-45	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3773-45C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3774-36	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3774-36C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3774-45	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3774-45C	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3780-28	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3781-28	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3781R-28ASP	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3782-28	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3782R-28ASP	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3783-28	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3783R-28ASP	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3783R-36ASP	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3784R-28ASP	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3786-28	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3810	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3810NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3815	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E3815NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4051-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

E4052-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4053-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4054-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4055-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4057-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4058-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4059-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4060-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4062-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E4073-CT	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7507	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7507-DB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7508	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7509	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7509-B	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7510-25	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7510-25DB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
E7512	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
ED20L	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ED21L	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EDW50	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EDW51	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EDW52	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEA21	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
EEA2135	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEA25	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEA2535	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEA28	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

EEA2835	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEA31	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEA33	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAORVIL21A	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
EEAORVIL25A	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
EEAXL21	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL2135	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL25	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL2535	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL28	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL2835	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL31	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EEAXL33	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA30AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA30AV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA30AVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA30CTAV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA30CTAVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45AV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45AVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45CTAMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45CTAV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA45CTAVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA60AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA60AVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA60AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

EGIA60CTAMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIA60CTAVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIATRS45AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIATRS45AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIATRS60AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIATRS60AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EGIAUSHORT	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
EGIAUSTND	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
EGIAUXL	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
EL20L	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL21L	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL22LN	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL23LN	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL-415	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL-430	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL-440	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL-450	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
EL-451	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ELW30	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ELW33	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
ELW50	G70776080009	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
FT0510	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
FT3000	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
FT3000DB	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
GC-121	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC-122	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC-123	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC123G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

GC-124	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC-321	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC-322	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC-323	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GC-33-M	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA10038L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA10038S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA10048L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA10048S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA6025L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA6025S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA6038L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA6038S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA6048L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA6048S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA60MTC	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA60MTS	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA60XTC	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA60XTS	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA8038L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA8038S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA8048L	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA8048S	G70776080040	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA80MTC	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA80MTS	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA80XTC	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GIA80XTS	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-120	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

GL-121	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-122	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-123	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-124	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-125	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-126	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-127	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-128	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-129	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-181	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-182	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-182-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-183	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-191	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-192	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-222	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-223	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-223-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-224	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-225	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-226	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-228	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-282	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-302	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-303	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-321	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-322	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-323	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

GL-324	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-32-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-332	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-333	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-33-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-34-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-44-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-45-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-46-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-47-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-522	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-523	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-524	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-61-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-62-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-63-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-64-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-66-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-67-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-68-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-69-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL72MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL79MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-881	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-882	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-884	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-885	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-889	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

GL-889-2	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-88G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-890	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLJ-50-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLJ-51-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GL-J53-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLJ-54-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLJ-56-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLJ-57-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLJ-58-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS-121	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS-122	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS-123	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS-321	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS-322	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS-323	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLS323GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GLT-69-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-341L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-343L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-344L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-496L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-497L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-563L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-573L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-632L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-634L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-806L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

GMM-807L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-873L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMM-951	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-540-MG	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-541-MG	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-563-MG	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-690L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-89L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-90L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-951	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GMMT-961	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
GN-283	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GN-284	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GNJ-283	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GP-282	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-30-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-33-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-34-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-43-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-44-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS451-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS452-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS453-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS46M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS47M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-62-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-63-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-64-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

GS-65-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-66-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-67-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-822	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-823	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-824	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS82G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-831	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-832	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-833	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-834	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS-835	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GS83G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ-33-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ-34-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ36M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ37M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ46M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ47M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ-63-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
GSJ-64-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
HEM3335	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
HEM3348	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
HP0204CVE	G2S0776080072	5/23/2024	TUV 0123	TUV 0123	12/31/2028	
L-0223K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1-	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-102	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-103	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

L-104	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-11	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-111	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-112	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L113	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-114	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-115	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-116	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-12	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-13	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-14	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-15	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1551-K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1562-K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1669-K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1670-K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1739-K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1740-K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1742K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1745K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1750K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1755K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1756K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1785K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1787K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1791K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1792K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-1796K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

L-1897K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2-	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-20	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-22	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-23	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-24	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-25	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2574K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-26	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-27	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2747K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2748K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2749K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2750K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2751K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2752K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2753K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-28	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-2800	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-3	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-32	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-33	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-34	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-35	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-36	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-38	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-39	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-4	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

L-40	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-402	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-403	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-404	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-405	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-406	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-41	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-42	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-43	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-4323K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-5	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-5118K	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-71	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-72	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-73	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-74	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-75	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
L-76	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LF0225G	G10887950028	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
LF1212	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1212A	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1823	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1837	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1844	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1923	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1930T	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1937	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF1944	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

LF2019	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF3225	G10887950028	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
LF3225C	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
LF4418	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF5637	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LF5644	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
LL-101	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-102	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-103	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-104	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-111	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-112	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL113	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-114	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-115	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-116	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-221	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-222	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-223	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-224	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-225	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-226	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-233	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-322	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LL-323	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LS-636	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LS-637	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LS-638	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

LS-639	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
LS-640	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
MS100703	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
MS100705	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
MS100708	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
MS101003	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
MS101005	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
MS101008	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
N-1603G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2500	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2510	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2512	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2513	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2514	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2530	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2531	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2532	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2533	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2540	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2541	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2543	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2544	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2546	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2547	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2548	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2704K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2705K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2714K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

N-2716K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2717-K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2718K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2719K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2721-K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2730-K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2734K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2736K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2756K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2757K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2760K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2761K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2766K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2770-K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2771-K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2781K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2786K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2787K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2799-K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2966K	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-2967	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-59	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-63	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
N-64	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
NONB11STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB12LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB12SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB12STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

NONB12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB15LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB15STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB5LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB5SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB5STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
NONB8STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OBTNONB8ST	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OBTONB11ST	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OBTONB12ST	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OMS-PDB1000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OMS-PDBS2	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OMS-T10SB	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OMS-XB1	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
OMS-XB2	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB11LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB11LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB11STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB11STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12STFCS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB15STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

ONB15STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB5LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB5SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB5STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
ONB5STF2C	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
P-2756K	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
P-2771K	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
P-2790G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
PB-6713K	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
PB-6723K	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RC11STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RC12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RC12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RC8STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RELTACK10R	G70776080076	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RELTACK3X10	G70776080076	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RELTACK4XDPT	G70776080076	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RELTACK5R	G70776080076	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RELTACK5RDPT	G70776080076	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RELTACK8RDPT	G70776080076	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
RNONB12LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RNONB12STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RNONB15LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RNONB15STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RNONB8LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RNONB8STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RONB11LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RONB11STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

RSEAL	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RUNVCA8LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
RUNVCA8STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
S100000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
S100700	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
S101000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
S110000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
S-1172	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1172-G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1173	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1174	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1272	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1274	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1373	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1732K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1733K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1735	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1746K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1750K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-176	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1765K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1766K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1768K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1769K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1780K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1783K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-1789K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-182	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

S-183	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-184	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-185	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-187	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-193	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-194	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-195	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-196	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-197	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-199	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-206	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-243	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-244	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-245	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-246	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-254	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-255	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-2780K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-2782K	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-282	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-303	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-304	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-305	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-316	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-317	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-318	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-346	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-403	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

S-404	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-405	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-605	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-606	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-607	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-608	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
S-610	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SBS-1880G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SBS-1883G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SBS-1884G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SBS-1928G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-3625G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-3626G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-3634	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC3634G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC542	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-543	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5587G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5588G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5590G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5616G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5617G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5618-G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5626	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5626G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5627G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5635G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5637-G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SC-5638-G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5640	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5642G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5643G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5679	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5686G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5688G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC5689G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC5690G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5779	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-5780G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-585	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-586	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC586G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-587	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-643	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-644	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-690G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-691	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-692	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-693	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-693G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SC-694G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SCBA	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SCD-2864G	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SCD-3049G	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SCD-3058G	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SCD-3059G	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SCDA13	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SCDA26	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SCDA39	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SCDA48	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SCGAA	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SDN-5666G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SDN-5691G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SDN-5696G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SDN-5943G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SEA3705	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEA3710	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEA3715	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEA3720	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEA3725	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEL7010	CE01948	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEP6000-	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEP6015	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEP6000NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SEP6015NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
SIG30AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG30AV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG30AVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG30CTAV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG30CTAVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG45AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG45CTAMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG45CTAV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG45CTAVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SIG60AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG60CTAMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIG60CTAVM	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGADAPTSHORT	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SIGADAPTSTND	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SIGADAPTXL	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SIGPHANDLE	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SIGPSHELL	G2S0776080072	5/23/2024	TUV 0123	TUV 0123	12/31/2028	
SIGSDL45CTVT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGSDS30CTV	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGSDS30CTVT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRS45AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRS45AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRS60AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRS60AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRSB45AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRSB45AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRSB60AMT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SIGTRSB60AXT	G70776080050	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1510-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1596	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1613	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1614	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1614-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1615	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1624-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1625	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL1625G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SL1625GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL1644G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1645G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1653	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1654	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1655G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1656	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1657	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1658	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1659	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1660	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1689	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-1689G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL1689GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-171	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-172	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-431	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-432	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5620	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5621	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5626	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5626-GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5627	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5627-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL5627GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5628	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5628G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL5628GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SL-5629	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5632	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5632G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5633	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5633G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5636	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5636-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5636GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5637	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5638	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5640	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5640G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5641	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5641G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5678	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5679	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5679G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5680	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5686	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5686G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5686-GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5687	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5687G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL5687GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5688	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5688-G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5690	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-5691G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SL5691GMDL	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-586	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-587	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-603	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-605	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-607	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-608	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-611	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-612	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-613	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-622	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-630	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-631	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-632	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-634	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-635	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-635G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-636	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-637	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-638	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-639	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-642	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-643	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-644	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-652	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-653	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-654	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-674	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SL-675	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-690	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-691	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-691G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-692	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-693	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-694	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-695	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-696	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-712	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-714	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-721	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-722	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-73-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-822	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-822G	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-823	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-83-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SL-85-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SMBTTOVLX	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SMBTTRNDX	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SMCYLCST	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SMM-5033	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5042	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5043	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5044	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5221	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5241	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SMM-5323	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5431	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5516	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5516G	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5526	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5536	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMM-5831	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
SMSBTOVLX	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SMSBTRNDX	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
SN-1647	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1647G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1689	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1689G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1693	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1954	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1955	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1956	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1964	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1966	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1994	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-1995	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-247	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-367	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-368	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-3686	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-3695	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-3695-GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-3697	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SN3697GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-3965	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-552G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5660G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5661G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5662	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5662G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5662GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5663	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5663G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5663GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5665G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5666	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5666G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5666GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5667	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5667G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5667GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5668	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5668G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5668GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5669	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5669G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5669GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5670	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5677	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-568	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5690	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SN-5690G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5691	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5694	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5694G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5696	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5696G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5696GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5697	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5698	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5698G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5698GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5699	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5699G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5699GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5767G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-5862G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN5862GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-615	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-616	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-617	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-626	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-627	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-628	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-629	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-630	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-631	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN641	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-643	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SN-644	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-648	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN648G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-649	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-653	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-658	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-659	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-659G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-660	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-661	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-661G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN661GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-662	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-662G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-663	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-663G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN663GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-664	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-664G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-665	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-666	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-667	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-667G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-669	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-670	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-671	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-671G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-672	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SN-673	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-673G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-674	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-681	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-682	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-688	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-689	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-693	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-694	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-728	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-747	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-750	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-758	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-760	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-761	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-762	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-763	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-764	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-764G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN764GMDL	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN765	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-770G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-771	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-771G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-772G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-775	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-776	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN776G	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SN-777	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-781	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-782	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-788	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-790	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-871	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-872	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SN-881	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1606	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1617	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1618	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1634	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1635	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1636	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1643	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1644	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP1647G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1693G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1696	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1696G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-1697	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-2698G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5563	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5587	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-559G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5663	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5663G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5680G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SP-5681	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5681G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5682	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5682G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5686	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5686G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5687	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5687G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5692G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5695	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5695-G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5697G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5698	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5698-G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5699	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-5699G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-621	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-622	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-622-BC	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP622G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-623	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-623-BC	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-624-BC	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-629	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-631	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-649	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-655	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-660	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SP-660G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-661	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-661G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-665	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-665G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-670	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-675	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-683	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-683G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-684	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-684G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-685	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-689	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-690	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP690G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-868	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SP-868G	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-1213G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-1233G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-1623G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-1643G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB5142G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-5143G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-5223G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-5413G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-5433G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-5633G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SPB-5833G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SPB-5843G	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-1639G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-1694G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-523	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-525	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5639	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5640	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5641	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5641G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5649G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5676	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5677	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5678	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5679	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5684	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-5685G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-621	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-622	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS622G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-623	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-624	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-629	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-632	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-633	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-645	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-646	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-647	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-648	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SS-651	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-652	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-653	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-654	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-655	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-656	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-673	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-675	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-677	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-678	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-679	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-680	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-681	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-682	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-683	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-683G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-684	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-684G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-685	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-685G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-686	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-689	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-694	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-695	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-70-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-722	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-723	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS723G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

SS-732	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-733	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-734	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-745	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-746	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-784	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-785	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-786	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-787	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
SS-82-M	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA3035L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA3035S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA3048L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA3048S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA30V3L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA30V3S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA4535L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA4535S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA4548L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA4548S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA6035L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA6035S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA6048L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA6048S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA9035L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA9035S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA9048L	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TA9048S	G70776080074	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

TRIEEA21MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA21XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA25MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA25XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA28MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA28XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA31MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA31XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA33MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEA33XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL21MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL21XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL25MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL25XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL28MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL28XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL31MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL31XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL33MT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
TRIEEAXL33XT	G10887950029	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-201	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-202	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-203	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC203G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-204	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-213	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC213G	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-214	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

UC-403	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-404	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-863	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-879	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UC-978	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-046	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-101	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-102	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-146	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-201	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-201-2	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-202	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-203	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-204	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-205	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-212	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-213	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-214	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-215	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-216	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-240	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-245	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-246	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-303	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-46-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-47-MG	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-676	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-876	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

UL-877	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-878	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-879	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UL-J-56-M	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
UNVCA11STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
UNVCA12LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
UNVCA12STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
UNVCA12STS	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
UNVCA5LGF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
UNVCA5SHF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
UNVCA5STF	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VCD-110	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2030556MFE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2030556MSE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20305FE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20305MFE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20305MSE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20305SE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20316FE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20316MFE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20316MSE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20316SE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2031MFE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2031MSE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD20331FE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2036305FE	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2202	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD2203	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VCD-256	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD-320	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD-346	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD-375	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD-656	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD8011G	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VCD8051G	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VL2610	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2610DB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2610E	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2610NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2615	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2615DB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2615E	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VL2615NSB	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VLFT10GEN	CE00500	5/26/2024	BSi 2797	BSi 2797	12/31/2028	
VLOCA004L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA006L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA008L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA204L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA206L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA208L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA304L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA306L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCA308L	G70776080085	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0003	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0004	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0013	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VLOCL0014	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0015	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0023	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0024	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0025	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0033	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0034	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0114	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0115	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0123	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0124	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0125	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0133	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0134	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0135	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0223	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0224	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0305	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0306	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0315	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0316	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0324	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0325	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0326	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0335	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0336	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0344	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0345	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VLOCL0346	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0416	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0426	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0436	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0603	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0604	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0613	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0614	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0615	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0624	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0625	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0643	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0644	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0803	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0804	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0813	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0814	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0824	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL0844	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL1413	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL1544	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL1545	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL1904	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2105	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2145	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2216	G70776080063	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2246	G70776080063	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2744	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VLOCL2815	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2825	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2826	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL2836	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL3216	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCL4636	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0003	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0004	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0005	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0013	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0014	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0015	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0023	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0024	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0025	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0033	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0034	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0035	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0113	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0114	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0115	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0123	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0124	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0125	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0133	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0134	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0135	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0223	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VLOCM0224	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0305	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0315	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0324	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0325	G70776080063	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0344	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0345	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0603	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0604	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0614	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0623	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0624	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0625	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0635	G70776080063	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0644	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0804	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0813	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0814	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0824	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0843	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM0844	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1203	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1204	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1413	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1423	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1544	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1545	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1624	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VLOCM1704	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1744	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1824	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1904	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM1944	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2004	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2044	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2105	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2115	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2145	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2205	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM2245	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM3225	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCM3245	G70776080063	5/2/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN004L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN006L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN008L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0305	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0306	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0325	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0326	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0327	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0346	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0604	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0605	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0614	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0615	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0644	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VLOCN0804	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN0814	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN1126	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN204L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN206L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN208L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN2146	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN2147	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN304L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN306L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VLOCN308L	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VMM-137	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
VP-102MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-120MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-126MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-151	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-204-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-205-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-209MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-331X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP331X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP363X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-365X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP365X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-37MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-421X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-422X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-435-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP-510X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-511-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-520X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-521-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-522-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-523	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-532X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-533	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-541X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-542-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-543	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP555X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP556MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP556X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP557MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP557X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP558X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-559-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP559X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP566X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP570X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP580X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP581X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP582X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP583X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP584X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP585X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-621-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP-628	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-630-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-631-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-652-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-653-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-660-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-681-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-701-MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP701X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP702MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP702X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP703MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP703X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-704-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-704-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP705MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP705X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP706MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP706X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-707-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-708-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP709MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP709X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-70-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-710-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-710-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-711-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-712-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP-712-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP713MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP713X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-714-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-715-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-718-MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP718X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-719-MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP719X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP720MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP720X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-721-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-721-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP725MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP725X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP726MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP726X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP727X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-728-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-728-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-72-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-72-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-733-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-733-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-734-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-735-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-735-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-736-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP-737-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-738-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-738-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-73-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-73-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-743-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-744-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-744-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-745-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-746-MX-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP746X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP746XG	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-747-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-747-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-74-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-751-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-753-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-754-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-755-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP757X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-759-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-75-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-760-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-761-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-761-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-762-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP763MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-763-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP-765-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-76-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-76-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-771-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-772-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-776-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-78-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP800X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-812-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP812X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP813X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-831-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-832-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-833	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-834	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-841-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-842-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-843	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-851-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-853	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-854-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP86X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP870X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP871X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP871XG	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP872X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-875-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-875-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP881X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP889X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP890X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-891	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-897-M	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-900-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-900-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-901-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-902-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-902-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-903-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-904-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-904-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-911-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-914-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-924-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-925-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-926-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-934-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-935-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-936-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-937	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-941-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-942-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-945	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-956	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-965	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP972X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VP-973-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-975-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-976-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-977	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-978	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VP-986-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-354-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-521-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-522-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-543	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-557-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-610-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-702-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-703-MX	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-703-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-704-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-706-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-709-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-710-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-711-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-713-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-720-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-726-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-728-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-733-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-735-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-776-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPF-807-X	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VPG36FE	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPG36SE	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPG56FE	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPG56SE	G70776080046	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VPP557X-2	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS070000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS100700	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS100705	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS100711P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS100712P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101005	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101011P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101012P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101015P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101500	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101505	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS101512P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS150000	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VS-533	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-552	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS581-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-706M-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS709-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-766G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS802-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS806-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS809-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

VS81G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS823-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-82-G	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-842	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-843	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-844	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-845	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-846	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-863	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS870-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS871-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS872-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS873-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS880-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS881-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS882-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS889-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS890-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS891-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-89-G-2	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VS-964	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
VSR100005	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
VSR100812P	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
WPLGR914	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
WPMD509	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
WPSM256	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
WPXLGR1117	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	
WPXSM24	G10776080079	5/26/2024	TUV 0123	TUV 0123	12/31/2028	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

X-1007	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1070	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1076	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1087	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1104	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1119	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1136	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1137	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1146	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1147	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1154	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1157	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1168	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1178	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1179	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1181	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1202	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1208	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1209	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1216	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1218	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1220	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1226	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1230	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1237	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1242	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1252	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1280	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

X-1302	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1307	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1313	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1317	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1319	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1323	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1325	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1326	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1327	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1331	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1353	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1354	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X1355	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X1356	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X-1357	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1360	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1362	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X-1367	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X1369	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X1370	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X1374	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X1378	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X1391	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
X1392	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X1394	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
X-7005	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2100	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2102	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

XX2103	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2104	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2106	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2107	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2108	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2109	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2110	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2112	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2114	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2116	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2118-L	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2119	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2120	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX2121-2	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2141	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2150	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2204	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2205	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-2206	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX2207	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX2210	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX2211	G70776080066	3/12/2024	TUV 0123	TUV 0123	12/31/2027	
XX2219	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-3300	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5000	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5011	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5027	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5056	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

XX-5058	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5059	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5060-L	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5069	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5072-L	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5087	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5090	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5092	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5095-L	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5099	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX5101	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5107	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5108	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5115	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5117	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5122	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5125	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5126	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5128	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5129	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5130	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5132	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5140	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5141	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5142	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5144	G70776080032	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5148	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5149	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

XX-5152	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5154	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5158	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5159	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5161	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5167	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5168	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5172	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX5183	G70776080078	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5185	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5186	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5190J	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5191J	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5192	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5193J	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5194J	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5195-J	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5202J	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5204J	G70776080082	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5205J	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5209	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5213	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5224	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5226	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5231	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5233	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5234	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5235	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	

Document No.:	RE00492884	Revision B
Document Title:	EU MDR Extension Manufacturer Declaration Letter	
Owner:	Regulatory	

XX-5239	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5241	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5242	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5246	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5247	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5249	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5253	G70776080030	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5255	G70776080086	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5259	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5263	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX5266	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5267	G70776080081	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5272	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5277	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5278	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5279	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5280	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5281	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5282	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5284	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5285	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	
XX-5286	G70776080027	5/26/2024	TUV 0123	TUV 0123	12/31/2027	



**Add value.
Inspire trust.**

TÜV SÜD Product Service GmbH · Ridlerstr. 65 · 80339 Munich · Germany

Covidien Ilc
15 Hampshire Street
Mansfield, MA 02048
USA

Your reference/letter of	Our reference/name	E-mail	Tel. extension	Date	Page
77608	713329955	[REDACTED]		2024-03-05	1 of 21

**TÜV SÜD Product Service GmbH
Confirmation Letter
CL 077608 0120 Rev. 00**

Reference: 713329955

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation EU 2023/607 amending Regulations (EU) 2017/745 (in the following referenced as MDR) as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices.

With this letter TÜV SÜD Product Service GmbH, designated under MDR and identified by the number 0123 on NANDO, confirms that we have received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the above stated manufacturer with the following SRN Number:

SRN Number: US-MF-000028763

The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below:

- Table 1 identifies the devices for which an MDR application has been received, written agreement concluded and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive.
- Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but TÜV SÜD Product Service GmbH has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

Registered Office: Munich
Trade Register Munich HRB 85 742
UniCredit Bank AG · BIC HYVEDEMMXXX
IBAN DE13 7002 0270 0048 8522 11
VAT ID No. DE129484267
Information pursuant to § 2 [1] DL-InfoV
(Germany) at tuvsud.com/imprint



TÜV SÜD Product Service GmbH
Application Review
Ridlerstr. 65
80339 Munich
Germany

tuvsud.com/ps
Hotline: +49 89 50084-747





If devices covered by certificates issued under Directive 90/385/EEC (AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that:

- the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry; or
- provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively.

The transition timelines in accordance Article 120 (3) of MDR that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120 (3c) of MDR, are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices (except sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition, measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

The issuance of the first confirmation letter is free of charge. We reserve the right to invoice further copies, amendments and / or changes of the confirmation letter according to effort.

For confirmation letter validity see: www.tuvsud.com/ps-cert?q=cert:CL 077608 0120 Rev. 00

In case of inquiries please contact:

On behalf of the Notified Body TÜV SÜD Product Service GmbH,
2024-03-05

[REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]
 [REDACTED]



Table 1: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 1 V-Loc™ PBT Non-absorbable Reload Basic UDI-DI: 0763000B000057688	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 2 Polysorb™ Coated Braided Synthetic Absorbable Sutures Basic UDI-DI: 0763000B00000977P	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0030 Rev. 01 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 3 Endo GIA™ Reloads with Tri-Staple™ Technology Endo GIA™ Gray Articulating Reloads Tri-Staple™ 2.0 Intelligent Reloads Basic UDI-DI: 0763000B00001106N	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0050 Rev. 03 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1)



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Endo GIA™ Reinforced Reloads with Tri-Staple™ Technology Tri-Staple™ 2.0 Reinforced Intelligent Reloads Basic UDI-DI: 0763000B000058487			Evidence #1; CA# Evidence #2; CA#
Device 4 EEA Staplers: EEA™ Auto Suture™ Circular Stapler with DST Series™ Technology Basic UDI-DI: 0763000B000011774 EEA™ Auto Suture™ Hemorrhoid and Prolapse Stapler with DST Series™ Technology Basic UDI-DI: 0763000B000011876 EEA™ Circular Stapler with Tri-Staple™ Technology Basic UDI-DI: 0763000B00012527J	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 5 Endo GIA™ Ultra Universal Staplers Basic UDI-DI: 0763000B00001116Q	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 6 Appose™ ULC Auto Suture™ Slim Body Skin Stapler 35 Basic UDI-DI: 0763000B00001126S Appose™ ULC Auto Suture™ Slim Body Skin Stapler 35W Basic UDI-DI: 0763000B000022374	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 7 V-loc™ 180 Absorbable Wound Closure Device Basic UDI-DI: 0763000B00001957Q V-loc™ 90 Absorbable Wound Closure Device Basic UDI-DI: 0763000B000020357F	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0063 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 8 Lapro-Clip™ Auto Suture™ Reusable Clip Applier Basic UDI-DI: 0763000B00003157A	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 9 Lapro-Clip™ Auto Suture™ Absorbable Ligating Clip Cartridge Basic UDI-DI: 0763000B00001907E	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0083 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 10 VersaOne™ Fascial Closure System Bladed Trocar with Fixation Cannula VersaOne™ Fascial Closure System Optical Trocar with Fixation Cannula Basic UDI-DI: 0763000B000021677	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 11 Maxon™ and Maxon™ CV Monofilament Absorbable Suture Basic UDI-DI: 0763000B000030577	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0066 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 12 GIA™ Auto Suture™ Stapler with DST Series™ Technology Basic UDI-DI: 0763000B00003077B GIA™ Auto Suture™ Loading Unit with DST Series™ Technology Basic UDI-DI: 0763000B000030679	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0040 Rev. 01 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 13 TA™ Auto Suture™ Stapler with DST Series™ Technology Basic UDI-DI: 0763000B00003097F TA™ Auto Suture™ Loading Unit with DST Series™ Technology Basic UDI-DI: 0763000B00003087D	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0074 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 14 V-Loc™ PBT Non-Absorbable Wound Closure Device Basic UDI-DI: 0763000B00001947N	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 15 Trocars (Thorocoport) Basic UDI-DI: 0763000B000057382	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 16 Signia Adapter Basic UDI-DI: 0763000B000010873	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 17 Trocars (VersaOne™) Basic UDI-DI: 0763000B000021677	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 18 Specimen Retrieval Bags: Endo Catch™ Gold Endo Catch™ II Basic UDI-DI: 0763000B00001917G Endo Bag™ Basic UDI-DI: 0763000B00001897V	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 19 Signia™ Power Handle Basic UDI-DI: 0763000B000010873	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 20 Surgiclip & Endo Clip: Premium Surgiclip™ Auto Suture™ Clip Applier Premium Surgiclip™ II Auto Suture™ Clip Applier Premium Surgiclip™ III Clip Applier Basic UDI-DI: 0763000B000031274 Endo Clip™ Auto Suture™ Clip Applier	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Endo Clip™ II Auto Suture™ Clip Applier Endo Clip™ III Auto Suture™ Clip Applier with Clip Logic™ Technology Basic UDI-DI: 0763000B000031274			
Device 21 Signia™ Power Shell Basic UDI-DI: 0763000B000055784	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G2S 077608 0072 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 22 Veriset™ Cutting Template Basic UDI-DI: 0763000B00001997Y	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G2S 077608 0072 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 23 Surgipro™ and Surgipro™ II Monofilament Polypropylene Basic UDI-DI: 0763000B00001967S 0763000B000026528D	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0081 Rev. 00 NB #: 0123 or



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
	<input type="checkbox"/> Class III implantable custom-made-device		<input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 24 Endo Hand™ Instruments: Endo Grasp™ Auto Suture™ Grasper Endo Clinch™ II Auto Suture™ Grasper Basic UDI-DI: 0763000B00002206W Endo Paddle Retract™ Auto Suture™ Paddle Retractor Endo Mini-Retract Endo Retract™ Auto Suture™ Fan Retractor Endo Retract™ II Auto Suture™ Articulating Fan Retractor Basic UDI-DI: 0763000B000022272 Endo Babcock™ Auto Suture™ Clamp Basic UDI-DI: 0763000B00002216Y Endo Peanut™ Auto Suture™ Blunt Dissector Basic UDI-DI: 0763000B00002197D	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 25 AbsorbaTack™ Fixation Device Basic UDI-DI: 0763000B000056685	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0073 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 26 Ticron™ Basic UDI-DI: 0763000B00005557Y 0763000B00024878N 0763000B00026608C Surgidac™ Basic UDI-DI: 0763000B00005547W 0763000B000054883	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0086 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 27 Monosof™ Monofilament Nylon Nonabsorbable Suture Basic UDI-DI: 0763000B00005477Z Surgilon™ Braided Nylon Nonabsorbable Suture Basic UDI-DI: 0763000B00005817Z	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0082 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1)



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Dermalon™ Monofilament Nylon Nonabsorbable Suture Basic UDI-DI: 0763000B00005898H			Evidence #1; CA# Evidence #2; CA#
Device 28 Sofsilik™ Coated Braided Silk Suture Basic UDI-DI: 0763000B000054985 0763000B00005537U	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0027 Rev. 01 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 29 Endo Hand™ Instruments: Endo Shears™ Auto Suture™ Dissector Basic UDI-DI: 0763000B00002197D Endo Shears™ Auto Suture™ Shears Endo Sciz™ Auto Suture™ Shears Endo Mini-Shears™ Auto Suture™ Shears Basic UDI-DI: 0763000B000021779	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 30 PDB™ Auto Suture™ Round Shape Balloon	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa	<input checked="" type="checkbox"/> N/A or	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<p>PDB™ Auto Suture™ Kidney Shape Balloon</p> <p>Basic UDI-DI: 0763000B00005637X</p> <p>Extra View™ Auto Suture™ Round Shape Balloon</p> <p>Extra View™ Auto Suture™ Oval Shape Balloon</p> <p>Basic UDI-DI: 0763000B00005647Z</p> <p>Auto Suture™ Structural Balloon Trocar</p> <p>Basic UDI-DI: 0763000B000055988</p>	<p><input type="checkbox"/> Class I devices in sterile condition</p> <p><input type="checkbox"/> Class I devices with measuring function</p> <p><input type="checkbox"/> Class III implantable custom-made-device</p>	<p><input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD</p> <p>Individual Article number:</p>	<p>or</p> <p><input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1)</p> <p>Evidence #1; CA#</p> <p>Evidence #2; CA#</p>
<p>Device 31</p> <p>Novafil™ Monofilament Polybutester</p> <p>Basic UDI-DI: 0763000B00003177E</p>	<p><input checked="" type="checkbox"/> Class III</p> <p><input type="checkbox"/> Class IIb implantable (non-exempted)</p> <p><input type="checkbox"/> Class IIb / Class IIb implantable (exempted)</p> <p><input type="checkbox"/> Class IIa</p> <p><input type="checkbox"/> Class I devices in sterile condition</p> <p><input type="checkbox"/> Class I devices with measuring function</p> <p><input type="checkbox"/> Class III implantable custom-made-device</p>	<p><input checked="" type="checkbox"/> N/A</p> <p>or</p> <p><input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD</p> <p>Individual Article number:</p>	<p><input checked="" type="checkbox"/> Certification as follows:</p> <p>Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0032 Rev. 01 NB #: 0123</p> <p>or</p> <p><input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1)</p> <p>Evidence #1; CA#</p> <p>Evidence #2; CA#</p>
<p>Device 32</p> <p>Polysorb™ Absorbable Single Stitch Reload</p> <p>Basic UDI-DI: 0763000B00005788C</p> <p>Surgidac™ Polyester Single Stitch Reload</p> <p>Basic UDI-DI: 0763000B000059082</p>	<p><input checked="" type="checkbox"/> Class III</p> <p><input type="checkbox"/> Class IIb implantable (non-exempted)</p> <p><input type="checkbox"/> Class IIb / Class IIb implantable (exempted)</p> <p><input type="checkbox"/> Class IIa</p> <p><input type="checkbox"/> Class I devices in sterile condition</p> <p><input type="checkbox"/> Class I devices with measuring function</p> <p><input type="checkbox"/> Class III implantable custom-made-device</p>	<p><input checked="" type="checkbox"/> N/A</p> <p>or</p> <p><input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD</p> <p>Individual Article number:</p>	<p><input checked="" type="checkbox"/> Certification as follows:</p> <p>Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0085 Rev. 00 NB #: 0123</p> <p>or</p> <p><input type="checkbox"/> Evidence that a competent authority of a Member</p>



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Sofsilik™ Single Stitch Reload Basic UDI-DI: 0763000B000059082 V-Loc™ 180 Absorbable Reload Basic UDI-DI: 0763000B00020367H			State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 33 ProTack™ Auto Suture™ Single Use Fixation Device Basic UDI-DI: 0763000B00005778A	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 34 EEA™ OrVil™ Auto Suture™ Transoral Circular Stapler Anvil with Advancing Proximal Guide Suture Basic UDI-DI: 0763000B00001156Y EEA™ Auto Suture™ Reusable Sizer Basic UDI-DI: 0763000B00001146W	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 35 Auto Suture™ Premium Extractor Basic UDI-DI: 0763000B000055886	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input checked="" type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G2S 077608 0072 Rev. 00 NB #: 0123 or



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
	measuring function <input type="checkbox"/> Class III implantable custom-made-device		<input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 36 Endo Stitch™ Auto Suture™ Suturing Device Basic UDI-DI: 0763000B00003106Y Endo Close™ Auto Suture™ Trocar Site Closure Device Basic UDI-DI: 0763000B000020347D	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 37 VersaOne Bladeless Reusable Positioning Trocar System: VersaOne™ Bladeless Obturator Basic UDI-DI: 0763000B000026738M VersaOne™ Optical Obturator Basic UDI-DI: 0763000B000026738M VersaOne™ Reusable Positioning Cannula Basic UDI-DI: 0763000B000026748P VersaOne™ Seal Basic UDI-DI: 0763000B000026738M	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 38 Spacemaker™ Pro Access and Dissector System Basic UDI-DI: 0763000B00005617T Endo-Lube™ Auto Suture™ Seal and Instrument Lubrication Kit Basic UDI-DI: 0763000B00005627V	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 39 Clearify™ Visualization System Basic UDI-DI: 0763000B000056583 Clearify™ Trocar Wipe Basic UDI-DI: 0763000B000056787 Fred™ 2 Anti-Fog Kit Basic UDI-DI: 0763000B000031478	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G2S 077608 0072 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 40 Endo Universal™ 65 Auto Suture™ Universal Hernia Stapler Basic UDI-DI: 0763000B000057484 Multifire Endo Hernia™ Auto Suture™ Loading Unit Basic UDI-DI: 0763000B00026658N	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
<p>Device 41</p> <p>Surgitie™ Auto Suture™ Ligating Loop</p> <p>Basic UDI-DI: 0763000B00025708A</p> <p>Surgitie™ Auto Suture™ Ligating Loop with Delivery System</p> <p>Basic UDI-DI: 0763000B00005717W 0763000B00005727Y</p> <p>Surgiwip™ Auto Suture™ Suture Ligature</p> <p>Basic UDI-DI: 0763000B00025728E</p> <p>Surgiwip™ Auto Suture™ Suture Ligature with Delivery System</p> <p>Basic UDI-DI: 0763000B00025478F 0763000B00025488H</p>	<p><input checked="" type="checkbox"/> Class III</p> <p><input type="checkbox"/> Class IIb implantable (non-exempted)</p> <p><input type="checkbox"/> Class IIb / Class IIb implantable (exempted)</p> <p><input type="checkbox"/> Class IIa</p> <p><input type="checkbox"/> Class I devices in sterile condition</p> <p><input type="checkbox"/> Class I devices with measuring function</p> <p><input type="checkbox"/> Class III implantable custom-made-device</p>	<p><input checked="" type="checkbox"/> N/A</p> <p>or</p> <p><input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD</p> <p>Individual Article number:</p>	<p><input checked="" type="checkbox"/> Certification as follows:</p> <p>Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0009 Rev. 01 NB #: 0123</p> <p>or</p> <p><input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#</p>
<p>Device 42</p> <p>VersaStep™ Auto Suture™ Insufflation Needle</p> <p>VersaStep™ Auto Suture™ Radially Expandable Sleeves</p> <p>VersaStep™ Auto Suture™ Dilator and Cannula with Radially Expandable Sleeves</p> <p>Basic UDI-DI: 0763000B00005878D</p> <p>VersaStep™ Plus Auto Suture™ Dilator and Cannula with Radially Expandable Sleeves</p> <p>Basic UDI-DI: 0763000B00005888F</p>	<p><input type="checkbox"/> Class III</p> <p><input type="checkbox"/> Class IIb implantable (non-exempted)</p> <p><input type="checkbox"/> Class IIb / Class IIb implantable (exempted)</p> <p><input checked="" type="checkbox"/> Class IIa</p> <p><input type="checkbox"/> Class I devices in sterile condition</p> <p><input type="checkbox"/> Class I devices with measuring function</p> <p><input type="checkbox"/> Class III implantable custom-made-device</p>	<p><input checked="" type="checkbox"/> N/A</p> <p>or</p> <p><input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD</p> <p>Individual Article number:</p>	<p><input checked="" type="checkbox"/> Certification as follows:</p> <p>Certificate #: G1 077608 0079 Rev. 00 NB #: 0123</p> <p>or</p> <p><input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#</p>



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
VersaStep™ Auto Suture™ Radially Expandable Sleeves Basic UDI-DI: 0763000B00005807X Mini Step™ Auto Suture™ Dilator and Cannula Basic UDI-DI: 0763000B00005868B			
Device 43 ReliaTack™ Articulating Reloadable Fixation Device Basic UDI-DI: 0763000B00020537H ReliaTack™ Reload with Standard Purchase Tacks Basic UDI-DI: 0763000B00020527F	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0076 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 44 PTFE Polymer Pledgets Basic UDI-DI: 0763000B00020507B	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0046 Rev. 01 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Device 45 Caprosyn™ Monofilament Absorbable Suture Basic UDI-DI: 0763000B00020547K	<input checked="" type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 G7 077608 0078 Rev. 00 NB #: 0123 or <input checked="" type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 46 SurgiSleeve Basic UDI-DI: 0763000B00020577R	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input checked="" type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#
Device 47 Stainless Steel Suture Basic UDI-DI: 0763000B000031376	<input type="checkbox"/> Class III <input type="checkbox"/> Class IIb implantable (non-exempted) <input checked="" type="checkbox"/> Class IIb / Class IIb implantable (exempted) <input type="checkbox"/> Class IIa <input type="checkbox"/> Class I devices in sterile condition <input type="checkbox"/> Class I devices with measuring function <input type="checkbox"/> Class III implantable custom-made-device	<input checked="" type="checkbox"/> N/A or <input type="checkbox"/> Identification of the corresponding device under MDD/AIMDD Individual Article number:	<input checked="" type="checkbox"/> Certification as follows: Certificate #: G1 077608 0079 Rev. 00 NB #: 0123 or <input type="checkbox"/> Evidence that a competent authority of a Member State had granted acc. MDR, Art.59 (1) or Art.97 (1) Evidence #1; CA# Evidence #2; CA#



Table 2: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under MDR application)	MDR Device classification (as proposed by the manufacturer and verified during application review)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Not applicable	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> N/A	<input checked="" type="checkbox"/> N/A

Confirmation Letter Version History

Date	TÜV SÜD Product Service GmbH internal reference traceable to each version of the letter	Action
2024-03-04	713329955	Initial issue
2024-03-05	713329955	Correction of Basic UDI-DIs for Device #37 and #40



EC-CERTIFICATE

(Full quality assurance system)



This is to certify that the company

Günter Bissinger Medizintechnik GmbH

Hans-Theisen-Str. 1
79331 Teningen
Germany

has implemented and maintains a full quality assurance system which applies to the products at every stage from design to final controls.

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of

Annex II – excluding Section 4 of Council Directive 93/42/EEC concerning medical devices

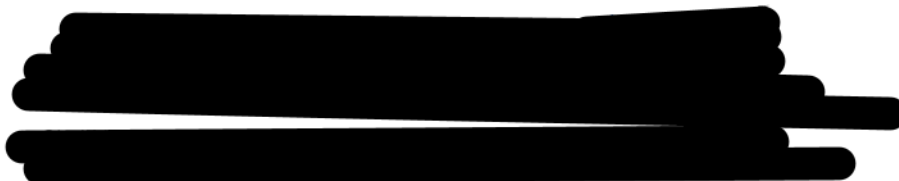
with respect to the following medical devices:

Non active instruments, RF surgical devices and accessories according to annex.

The manufacturer is subject to surveillance according to Annex II, Section 5. The CE marking with the Notified Body Identification Number (0297) may be affixed on the devices listed in the certificate. An EC Design Examination Certificate according to Annex II, Section 4 is required for class III devices covered by this certificate. The certificate is in the case of class I(s) devices (I(s) = class I products placed on the market in sterile conditions) limited to the aspects of manufacture concerned with securing and maintaining sterile conditions. The certificate is in the case of class I(m) devices (I(m) = class I devices with a measuring function) limited to the aspects of manufacture concerned with the conformity of the products with the metrological requirements.

Certificate registration no.	003171 MR2
Certificate unique ID	170775967
Effective date	2021-05-07
Expiry date	2024-05-26
Frankfurt am Main	2021-05-07

DQS Medizinprodukte GmbH



August-Schanz-Straße 21, 60433 Frankfurt am Main,

DQS Medizinprodukte GmbH is a Notified Body according to Council Directive 93/42/EEC concerning medical devices with the Identification Number 0297.



Annex to certificate

Certificate registration No.: 003171 MR2

Certificate unique ID: 170775967

Effective date: 2021-05-07

Günter Bissinger Medizintechnik GmbH

Hans-Theisen-Str. 1

79331 Teningen

Germany

Device family	Device	Class
RF surgical devices and accessories:	RF-surgery device Compact Coagulator 8070	IIb
	Bipolar instruments and accessories for RF-surgery	
	Monopolar instruments and accessories for RF-surgery	
	Monopolar electrodes in sterile and non-sterile version	
Non active instruments:	Bipolar forceps in sterile and non-sterile version	
	Uterine manipulator	IIa
	Trokar	IIa

Günter Bissinger Medizintechnik GmbH

Hans-Theisen-Str. 1
79331 Teningen
Germany

Date: 2024.04.16

Notified Body Confirmation Letter

Reference: 1000140188

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation EU 2023/607 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices

This letter confirms that, DQS Medizinprodukte GmbH, a Notified Body designated against Regulation (EU) 2017/745 (MDR) and identified by the number 0297 on NANDO, has received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the following manufacturer:

Günter Bissinger Medizintechnik GmbH

Hans-Theisen-Str. 1
79331 Teningen
Germany

SRN: (DE-MF-000005545)

The devices covered by the formal application and the written agreement mentioned above are identified in the Tables listed below: Table 1 identifies the devices for which an MDR application has been received, written agreement concluded and for which DQS Medizinprodukte GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive. Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but DQS Medizinprodukte GmbH has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

In the case of devices covered by certificates issued under Directive 90/385/EEC (AIMDD) or Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that the manufacturer signed the written agreement under MDR by the date of MDD/AIMDD certificate expiry, or provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively, by the 20 Mar 2023 for the relevant devices. The transition timelines that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 120.3c of MDR (as amended by (EU) 2023/607), are shown below:

- 26 May 2026 for Class III custom-made implantable devices
- 31 December 2027 for Class III devices and Class IIb implantable devices excluding Well-established technologies (WET - sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition or have a measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR (e.g., class I devices that qualify as re-usable surgical instruments)

[REDACTED]

Table 1: Devices covered by this letter and for which the NB is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name and Basic UDI-DI (as proposed by the manufacturer within the application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
POWERGRIP POWERGRIP/ORBITARIS ORBITARIS POWERGRIP 3.0 Monopolare Koagulationszange POWEREDGE CLASSIC Bipolare Elektrode für die MIC (5 mm) SLIMLINE Bipolare Elektrode für die MIC Slimline (3 mm) Bipolare Tastelektrode Bipolare Stichkoagulationselektrode Monopolare Elektrode für die MIC Monopolare Mikronadelelektrode Hysterektomieinstrument mit Drahtschlinge Monopolare Elektrode für die MIC (3 mm) Mikrodissektionsnadelelektrode 4250418720016H	Class IIb	HF-Chirurgiegeräte und Zubehör: - Monopolare Instrumente und Zubehör für die Elektrochirurgie - Bipolare Instrumente und Zubehör für die Elektrochirurgie - Monopolare Elektroden in steriler und unsteriler Ausführung	003171 MR2 ID# 170775967 NB 0297
HF-Handgriff Monopolare Elektroden 4250418720026K	Class IIb	HF-Chirurgiegeräte und Zubehör: - Monopolare Elektroden in steriler und unsteriler Ausführung - Monopolare Instrumente und Zubehör für die Elektrochirurgie	003171 MR2 ID# 170775967 NB 0297

Device name and Basic UDI-DI (as proposed by the manufacturer within the application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
BiTech Bipolare Scheren BlackLine bipolare Scheren 4250418720046P	Class IIb	HF-Chirurgiegeräte und Zubehör: - Bipolare Instrumente und Zubehör für die Elektrochirurgie	003171 MR2 ID# 170775967 NB 0297
Sterile Klinge für Poweredge Sterile Mikrodissektionsnadelelektroden 4250418720066T	Class IIb	HF-Chirurgiegeräte und Zubehör: - Bipolare Pinzetten in steriler und unsteriler Ausführung - Monopolare Elektroden in steriler und unsteriler Ausführung	003171 MR2 ID# 170775967 NB 0297
Bipolare Pinzette MICRODOME MITHRAS Monopolare Pinzette Zubehör Bipolare Pinzette 4250418720076V	Class IIb	HF-Chirurgiegeräte und Zubehör: - Bipolare Pinzetten in steriler und unsteriler Ausführung - Monopolare Instrumente und Zubehör für die Elektrochirurgie - Bipolare Instrumente und Zubehör für die Elektrochirurgie	003171 MR2 ID# 170775967 NB 0297
Monopolares Resektoskop PLASMALOOP 4250418720086X	Class IIb	HF-Chirurgiegeräte und Zubehör: - Bipolare Instrumente und Zubehör für die Elektrochirurgie - Monopolare Instrumente und Zubehör für die Elektrochirurgie	003171 MR2 ID# 170775967 NB 0297
Mangeshikar Uterus Manipulator Mangeshikar Uterus Manipulator Advanced + 4250418720096Z	Class IIa	Nichtaktive Instrumente: - Uterusmanipulator	003171 MR2 ID# 170775967 NB 0297

Table 2: Devices covered by this letter and for which the NB is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name and Basic UDI-DI (as proposed by the manufacturer within the application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD/AIMDD device	MDD/AIMDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Nadelhalter 4250418720126N	Class I devices that qualify as re-usable surgical instruments	N/A	N/A - Class I device under MDD (no NB required)

Confirmation Letter Revision History

Date	NB internal reference traceable to each version of the letter	Action
2024-04-16	1000140188	Initial issue
	Cert-ID	description of change (e.g. addition of device XYZ to Table 1)
	Cert-ID	description of change (e.g. removal of device XYZ from Table 2)

EC Certificate - Full Quality Assurance System

Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No.

CE 01906

Issued To:

**Fiab SpA
Via P. Costoli, 4
Vicchio
Firenze
50039
Italy**

In respect of:

The design, development and manufacture of sterile leads for transoesophageal cardiac and temperature monitoring, cardiac stimulation, cardiac defibrillation and electrophysiological studies; percutaneous introducers; electronic equipments for oesophageal temperature monitoring, electrophysiological studies and emergency cardiac stimulation; sterile and non sterile electrosurgical electrodes and related accessories; electrodes for defibrillation/pacing; sterile single use neuropacers; sterile single use and reusable electrocauteries and associated accessories; sterile and non sterile, single use and reusable needle electrodes for EEG and EMG.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 0086):

[Redacted Signature]

First Issued: **1998-05-11**

Date: **2018-05-10**

Expiry Date: **2023-05-10**

...making excellence a habit.™

Page 1 of 1

Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.
This certificate was issued electronically and is bound by the conditions of the contract.

BSI

CE SERTIFIKATAS - Visiškas Kokybės Užtikrinimas

Europos Tarybos Direktyva 93/42/EEC, Priedas II, Dalis 4

Nr. CE 01906

Kam išduota:

FIAB SpA
Via P. Costoli, 4
Vicchio
Firenze
50039
Italija

Dėl:

Kūrimo, projektavimo ir gamybos sterilių zondų perstempliniams širdies stebėjimams, stimuliacijai ir defibriliacijai; prietaisų elektrofiziologiniams tyrimams ir ekstrinei širdies stimuliacijai; sterilių ir nesterilių kaniulių, kaukių, rinkiniams ir priemonių deguonies terapijai, sterilių ir nesterilių elektrochirurginių rankenų antgaliai, elektrodai ir susijusių priemonių, sterilių vienkartinių neurostimuliatorių ir elektrokauterių

Remiantis mūsų atliktais patikrinimais pagal Europos Tarybos Direktyvą 93/42/EEC, Priedą II, Dalį 4.

Britų Standartų Institutas - įgaliota ir Notifikuotoji įstaiga aukščiau minėtai Direktyvai (Notifikuotos įstaigos numeris 0086):

(parašas)



Pirmoji sertifikavimo data: 1998 gegužės 11d.

Šio sertifikato data: 2018 gegužės 10 d.

Galioja iki: 2023 gegužės 10d.

(rekvizitai)

Vicchio (FI), 12/04/2023

TO WHOM IT MAY CONCERN

Subject: Extension of the MDR 2017/745 transitional period – confirmation of validity of FIAB MDD 93/42/EEC Certificates CE 01906, CE 649635, CE 720326

The amendment of the Medical Devices regulation (MDR) 2017/745 introduced by the *Regulation (EU) 2023/607 of the European Parliament and of the Council of 15 March 2023 amending Regulations (EU) 2017/745 (MDR) and (EU) 2017/746 (IVDR) as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices (Attachment 1 of this letter)* aims – among other things – to give Manufacturers and Notified Bodies sufficiently more time to carry out, in accordance with the MDR, the conformity assessment of devices covered by a certificate issued in accordance with Medical Devices Directive (MDD) 93/42/EEC that is going to expire or is already expired.

Such devices, also known as ‘legacy devices’ can benefit from an extended transitional period as set in the Regulation (EU) 2023/607, for the application of MDR.

‘Legacy devices’ should be understood as devices, which, in accordance with the MDR’s transitional provisions, are placed on the market after the MDR’s date of application (i.e. 26 May 2021) if certain conditions are fulfilled. Those devices covered by a valid EC certificate issued in accordance with MDD prior to 26 May 2021 benefit of an extension of the transitional period beyond 26 May 2024 if the conditions laid down in Article 120(3c) MDR are fulfilled, for the relevant certificates expired or going to expire after 20 March 2023.

As the Manufacturer of the medical devices listed in **Attachment 2** of this letter, FIAB SpA herewith confirms that the products covered by the following MDD 93/42/EEC certificates

- CE 01906, MDD Annex II.3 (Full Quality Assurance system certificate)
- CE 649635, CE 720326 MDD Annex II.4 (Design Dossier Examination certificate)

fulfil the requirements defined by Regulation (EU) 2023/607.

Consequently, the above mentioned certificates can be considered as valid, respectively, until 31/12/2028 for class IIa and class IIb medical devices (CE 01906) and until 31/12/2027 for class III medical devices (CE 649635, CE 720326), when FIAB SpA continues to comply with the relevant requirements in Regulation (EU) 2017/745 as amended by Regulation (EU) 2023/607.

The confirmation is made taking into account the following aspects

- Regulation (EU) 2023/607 extends the validity of CE certificates under MDD, considering limited capacity of Notified Bodies accredited for conformity assessment procedures under MDR
- Important condition of this extension is that the Manufacturer shall submit an MDR certification application for these devices to a MDR Notified Body not later than 26/05/2024 and shall sign MDR certification agreement with the MDR Notified Body no later than 26/09/2024
- Other requirements for this extension includes e.g.: the devices continue to comply with MDD there are no significant changes in the design and intended purpose; devices do not present an unacceptable risk to the health or safety; the Manufacturer has put in place a quality management system in accordance with MDR; a Notified Body is still performing surveillance activity



FIAB SpA is providing appropriate evidences demonstrating that the relevant requirements in Regulation (EU) 2017/745 as amended by Regulation (EU) 2023/607 have been fulfilled by now. In particular

- for each of the medical devices listed in **Attachment 2** of this letter, an MDR certification application was already submitted by FIAB to the MDR Notified Body 2797 (BSI) and the respective MDR certification agreement has been signed, as listed in Attachment 2;
- the devices continue to comply with MDD, according to the surveillance activity performed by the same Notified Body 2797 to FIAB; this ensures that there are no significant changes and the devices do not present an unacceptable risk;
- FIAB has already put in place a quality management system in accordance with MDR, as attested by the EU Quality Management System Certificate, MDR 747884 in **Attachment 3**, according to MDR Annex IX chapter I and III. Such MDR certificate already cover the medical devices for which the Notified Body 2797 completed the certification assessment

[REDACTED]
[REDACTED]
[REDACTED]
FIAB S.p.A. [REDACTED]
[REDACTED]
[REDACTED]

Vicchio (FI), 2023 04 12

VISIEMS, KAM TAI GALI BŪTI AKTUALU

Tema: MDR 2017/745 pereinamojo laikotarpio pratęsimas - FIAB MDD 93/42/EEB sertifikatų CE 01906, CE 649635, CE 720326 galiojimo patvirtinimas

Medicinos prietaisų reglamento (MDR) 2017/745 pakeitimu, padarytu 2023 m. kovo 15 d. Europos Parlamento ir Tarybos reglamentu (ES) 2023/607, kuriuo iš dalies keičiamos reglamentų (ES) 2017/745 (MDR) ir (ES) 2017/746 (IVDR) nuostatos dėl pereinamojo laikotarpio nuostatų, susijusių su tam tikrais medicinos prietaisais ir in vitro diagnostikos medicinos prietaisais (šio laiško **1 priedas**), siekiama, be kita ko, suteikti gamintojams ir notifikuotosioms įstaigoms pakankamai daugiau laiko, prietaisų, kuriems taikomas pagal Medicinos prietaisų direktyvą (MDD) 93/42/EEB išduotas sertifikatas, kurio galiojimas baigsis arba jau baigėsi, atitikties vertinimą pagal MDR.

Tokiems prietaisams, dar vadinamiems "senaisiais prietaisais", gali būti taikomas Reglamente (ES) 2023/607 nustatytas ilgesnis pereinamasis laikotarpis MDR taikyti.

"Senosios priemonės" turėtų būti suprantamos kaip priemonės, kurios, vadovaujantis MDR pereinamojo laikotarpio nuostatomis, pateikiamos rinkai po MDR taikymo pradžios datos (t. y. 2021 m. gegužės 26 d.), jei įvykdomos tam tikros sąlygos. Tiems prietaisams, kuriems taikomas galiojantis EB sertifikatas, išduotas pagal MDD iki 2021 m. gegužės 26 d., pereinamasis laikotarpis gali būti pratęstas po 2024 m. gegužės 26 d., jei įvykdomos MDR 120 straipsnio 3c dalyje nustatytos sąlygos dėl atitinkamų sertifikatų, kurių galiojimo laikas baigėsi arba baigsis po 2023 m. kovo 20 d.

Kaip šio laiško **2 priede** išvardytų medicinos prietaisų gamintojas, FIAB SpA patvirtina, kad gaminiai, kuriems išduoti šie MDD 93/42/EEB sertifikatai

- CE 01906, MDD II.3 priedas (Visiško kokybės užtikrinimo sistemos sertifikatas)
- CE 649635, CE 720326 MDD II.4 priedas (Projekto dokumentų rinkinio tyrimo

sertifikatas) atitinka Reglamente (ES) 2023/607 nustatytus reikalavimus.

Todėl pirmiau minėti sertifikatai gali būti laikomi galiojančiais atitinkamai iki 2028 m. gruodžio 31 d. IIa ir IIb klasės medicinos prietaisams (CE 01906) ir iki 2027 m. gruodžio 31 d. III klasės medicinos prietaisams (CE 649635, CE 720326), kai "FIAB SpA" ir toliau atitinka atitinkamus Reglamento (ES) 2017/745 su pakeitimais, padarytais Reglamentu (ES) 2023/607, reikalavimus.

Patvirtinimas atliekamas atsižvelgiant į šiuos aspektus

- Reglamentu (ES) Nr. 2023/607 pratęsimas CE sertifikatų galiojimas pagal MDD, atsižvelgiant į ribotus paskelbtųjų įstaigų, akredituotų atitikties vertinimo procedūroms pagal MDR, pajėgumus.
- Svarbi šio pratęsimo sąlyga yra ta, kad gamintojas ne vėliau kaip 2024-05-26 turi pateikti MDR notifikuotajai įstaigai šių prietaisų MDR sertifikavimo paraišką ir ne vėliau kaip 2024-09-26 turi pasirašyti MDR sertifikavimo sutartį su MDR notifikuotąja įstaiga.
- Kiti reikalavimai šiam pratęsimui: prietaisai ir toliau atitinka MDD, nėra reikšmingų konstrukcijos ir paskirties pakeitimų, prietaisai nekelia nepriimtino pavojaus sveikatai ar saugai, gamintojas įdiegė kokybės valdymo sistemą pagal MDR, notifikuotoji įstaiga vis dar atlieka priežiūros veikla.

"FIAB SpA" pateikia tinkamus įrodymus, kad atitinkami Reglamento (ES) 2017/745 su pakeitimais, padarytais Reglamentu (ES) 2023/607, reikalavimai jau įvykdyti. Visų pirma

- FIAB jau pateikė paraišką dėl kiekvieno iš šio rašto **2 priede** išvardytų medicinos prietaisų MDR sertifikavimo 2797 notifikuotajai įstaigai (BSI) ir pasirašė atitinkamą MDR sertifikavimo sutartį, kaip nurodyta 2 priede;
- prietaisai ir toliau atitinka MDD pagal tos pačios notifikuotosios įstaigos 2797 FIAB atliktą priežiūros veiklą; taip užtikrinama, kad nėra reikšmingų pokyčių ir prietaisai nekelia nepriimtinos rizikos;
- FIAB jau yra įdiegusi kokybės valdymo sistemą pagal MDR, tai patvirtina ES kokybės valdymo sistemos sertifikatas, MDR 747884, pateiktas **3 priede**, pagal MDR IX priedo I ir III skyrius. Toks MDR sertifikatas jau taikomas medicinos prietaisams, kurių sertifikavimo vertinimą atliko notifikuotoji įstaiga 2797

[redacted]
[redacted]
[redacted]
S.p.A.

el. paštas
[redacted]
[redacted]

LSMU ligoninei Kauno klinikos VŠĮ

TIEKĖJO PATVIRTINIMAS

2025 01 20

Kaunas

UAB „Sorimpeksas“ patvirtiname:

- visi priedai techniškai suderinami su LSMU ligoninėje Kauno klinikose naudojamais elektrochirurginiais prietaisais BOWA ARC 350;
- garantinis terminas 12 mėnesių (reikalavimas taikomas daugkartinio naudojimo priedams).