	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
	I. General Requirements		<b>,</b>		
1	Devices shall achieve the performance intended by the manufacturer and be designed and manufactured in such a way that, during normal conditions of use, they are suitable for their intended purpose, taking into account the generally acknowledged state of the art. They shall not compromise the clinical condition or the safety of patients, or the safety and health of users or, where applicable, other persons, provided that any risks which may be associated with their use constitute acceptable risks when weighed against the benefits to the patient and are compatible with a high level of protection of health and safety. This shall include: – reducing as far as possible the risk of use error due to ergonomic features of the device and the environment in which the device is intended to be used (design for				
-	patient safety), and – consideration of the technical knowledge, experience, education and training, and the medical and physical conditions of intended users (design for lay, professional, disabled or other users).				
2	The solutions adopted by the manufacturer for the design and manufacture of the devices shall conform to safety principles, taking account of the generally acknowledged state of the art. To reduce risks, the manufacturer shall manage the risks so that the residual risk associated with each hazard as well as the overall residual risk is judged acceptable. The manufacturer shall apply the following principles in the priority order listed: (a) identify known or foreseeable hazards and estimate the associated risks arising from the intended use and foreseeable misuse;				
	<ul> <li>(b) eliminate risks as far as possible through inherently safe design and manufacture;</li> <li>(c) reduce as far as possible the remaining risks by taking adequate protection measures, including alarms; and</li> <li>(d) provide training to upper and/or inform upper of any residual risks</li> </ul>				
3	The characteristics and performances of the device shall not be adversely affected to such a degree that the health or safety of the patient or the user and, where applicable, of other persons are compromised during the lifetime of the device, as indicated by the manufacturer, when the device is subjected to the stresses which can occur during normal conditions of use and has been properly maintained in accordance with the manufacturer's instructions. When no lifetime is stated, the same applies for the lifetime reasonably to be expected of a device of that kind, having				





	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
	regard to the intended purpose and the anticipated use of the device.				
4	Devices shall be designed, manufactured and packaged in such a way that their				
	characteristics and performances during their intended use will not be adversely				
	affected by transport and storage conditions (for example, fluctuations of temperature				
	and numberly) taking account of the instructions and information provided by the manufacturer				
5	All known and foreseeable risks, and any undesirable side-effects, shall be minimized				
Ŭ	and be acceptable when weighed against the benefits to the patient of the achieved				
	performance of the device during normal conditions of use.				
6	For devices listed in Annex XV for which the manufacturer does not claim a medical				
	purpose, the general requirements set out in Sections 1 and 5 shall be understood				
	that the device, when used under the conditions and for the purposes intended, shall				
	not present any risk or only the minimum acceptable risks related to the product's use				
	which is consistent with a high level of protection for the safety and health of persons.				
II. Requirements Regarding Design and Co					
7	Chemical, physical and biological properties	ſ			
1.1	The devices shall be designed and manufactured in such a way as to ensure the abaracteristics and performance referred to in Chapter L'Constral Requirements'				
	Particular attention shall be naid to:				
	(a) the choice of materials used particularly as regards toxicity and where				
	appropriate. flammability:				
	(b) the compatibility between the materials used and biological tissues, cells, and				
	body fluids taking account of the intended purpose of the device;				
	(c) where appropriate, the results of biophysical or modeling research whose validity				
	has been demonstrated beforehand;				
	(d) the choice of materials used, reflecting, where appropriate, matters such as				
7.0	hardness, wear and fatigue strength.				
1.2	The devices shall be designed, manufactured and packaged in such a way as to				
	the intended purpose of the device, and to the persons involved in the transport				
	storage and use of the devices. Particular attention shall be haid to tissues exposed				
	and to the duration and frequency of exposure.				
7.3	The devices shall be designed and manufactured in such a way that they can be				
	used safely with the materials and substances, including gases, with which they enter				
	into contact during their normal use or during routine procedures; if the devices are				



Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
intended to administer medicinal products they shall be designed and manufactured in such a way as to be compatible with the medicinal products concerned according to the provisions and restrictions governing these medicinal products and that both the performance of the medicinal products and of the devices are maintained in accordance with their respective indications and intended use.				
<b>7.4</b> The devices shall be designed and manufactured in such a way as to reduce as far as possible and appropriate the risks posed by substances that may leach or leak from the device. Special attention shall be given to substances which are carcinogenic, mutagenic or toxic to reproduction, in accordance with Part 3 of Annex VI to Regulation (EC) No 1272/2008 of the European Parliament and of the Council of 16 December 2008 on classification, labeling and packaging of substances and mixtures, amending and repealing Directives 67/548/EEC and 1999/45/EC, and amending Regulation (EC) No 1907/2006 (OJ L 353, 31.12.2008, p.1), and to substances having endocrine disrupting properties for which there is scientific evidence of probable serious effects to human health and which are identified in accordance with the procedure set out in Article 59 of Regulation (EC) No 1907/2006 of the European Parliament and of the Council of 18 December 2006 concerning the Registration, Evaluation, Authorization and Restriction of Chemicals (REACH) (OJ L 136, 29.5.2007, p.3).				
If devices, or parts thereof, that are intended – to be invasive devices and to come into contact with the body of the patient for short- or long-term, or – to (re)administer medicines, body liquids or other substances, including gases, to/from the body, or – to transport or store such medicines, body fluids or substances, including gases, to be (re)administered to the body				
<ul> <li>contain, in a concentration of 0.1% by mass of the plasticized material or above, phthalates which are classified as carcinogenic, mutagenic or toxic to reproduction of category 1A or 1B in accordance with Part 3 of Annex VI to Regulation (EC) No 1272/2008, these devices shall be labeled on the device itself and/or on the packaging for each unit or, where appropriate, on the sales packaging as devices containing phthalates.</li> <li>If the intended use of such devices includes treatment of children or treatment of</li> </ul>				



		Applicable to	Method Used to	Method	Reference to Supporting
	Essential Requirements	the Device?	Demonstrate Conformity	Reference	Controlled Documents
	pregnant or nursing women, the manufacturer shall provide a specific justification for				
	the use of these substances with regard to compliance with the general safety and				
	performance requirements, in particular of this paragraph, within the technical				
	documentation and, within the instructions for use, information on residual risks for				
7.5	these patient groups and, if applicable, on appropriate precautionary measures.				
1.5	Devices shall be designed and manufactured in such a way as to reduce as far as				
	possible and appropriate risks posed by the uninternitorial ingress of egress of				
	the environment in which it is intended to be used				
76	The devices shall be designed and manufactured in such a way as to reduce to a				
7.0	minimum the risks linked to the size and the properties of particles used. Special care				
	shall be applied when devices contain or consist of particles discu. Opeolar care				
	into the patient's or user's body.				
8	Infection and microbial contamination				
8.1	The devices and manufacturing processes shall be designed in such a way as to				
	eliminate or to reduce as far as possible the risk of infection to patients, users and,				
	where applicable, other persons. The design shall:				
	(a) allow easy handling,				
	and, where necessary,				
	(b) reduce as far as possible and appropriate any microbial leakage from the device				
	and/or microbial exposure during use,				
	(c) prevent microbial contamination of the device or specimen.				
8.2	Devices labeled as having a special microbiological state shall be designed,				
	manufactured and packaged to ensure that they remain so when placed on the				
	market and remain so under the transport and storage conditions specified by the				
83	Devices delivered in a sterile state shall be designed, manufactured and packaged in				
0.5	a pon reusable pack, and/or according to appropriate procedures, to ensure that they				
	are sterile when placed on the market and remain sterile under the transport and				
	storage conditions indicated by the manufacturer until the protective packaging is				
	damaged or opened.				
8.4	Devices labeled either as sterile or as having a special microbiological state shall				
	have been processed, manufactured and, if applicable, sterilized by appropriate,				
	validated methods.				
8.5	Devices intended to be sterilized shall be manufactured in appropriately controlled				

Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking Identity of the device and applicable configurations/variants covered by this checklist:

	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate	Method Reference	Reference to Supporting Controlled
	(a.g. anviranmental) conditions		Conformity		Documents
9.6	Ce.g. environmental) conditions.				
0.0	of the product and if the dovices are to be sterilized prior to use, minimize the risk of				
	microbial contamination: the packaging system shall be suitable taking account of the				
	method of sterilization indicated by the manufacturer				
87	The labeling of the device shall distinguish between identical or similar products				
0.7	placed on the market in both sterile and non-sterile condition				
9	Devices incorporating a substance considered to be a medicinal product and devintended to be ingested, inhaled or administered rectally or vaginally	vices composed o	of substances or	combination c	f substances
9.1	In the case of devices referred to in the first subparagraph of Article 1(4), the quality,				
	safety and usefulness of the substance which, if used separately, would be				
	considered to be a medicinal product within the meaning of Article 1 of Directive				
	2001/83/EC, shall be verified by analogy with the methods specified in Annex I to				
	Directive 2001/83/EC, as laid down in the applicable conformity assessment				
	procedure in this Regulation.				
9.2	Devices that are composed of substances or combination of substances intended to				
	be ingested, innaled or administered rectally or vaginally and that are absorbed by or				
	requirements loid down in Anney Lto Directive 2001/82/EC				
10	Devices incorporating materials of biological origin				
10 1	Ear devices manufactured utilizing tissues or cells, or their derivatives, of human				
10.1	origin which are covered by this Regulation in accordance with point (e) of Article				
	1(2) the following applies:				
	(a) Donation procurement and testing of tissues and cells of human origin used for				
	the manufacture of devices shall be made in accordance with Directive 2004/23/EC				
	(b) The processing, preservation and any other handling of those tissues and cells				
	shall be carried out so as to provide optimal safety for patients, users and, where				
	applicable, other persons. In particular, safety with regard to viruses and other				
	transmissible agents shall be addressed by implementation of validated methods of				
	elimination or inactivation in the course of the manufacturing process.				
	(c) It shall be ensured that the traceability system for devices manufactured utilizing				
	those human tissues or cells is complementary and compatible with the traceability				
	and data protection requirements laid down in Directive 2004/23/EC and in Directive				
	2002/98/EC of the European Parliament and of the Council of 27 January 2003				
	setting standards of quality and safety for the collection, testing, processing, storage				

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	Essential Requirements	Applicable to	Method Used to	Method	Reference to Supporting
		the Device?	Demonstrate Conformity	Reference	Controlled Documents
	and distribution of human blood and blood components and amending Directive		2		
10.2	For devices manufactured utilizing tissues or cells, or their derivatives, of animal				
	origin which are non-viable or rendered non-viable the following applies:				
	(a) Where feasible taking into account the animal species, tissues and cells of animal				
	origin shall originate from animals that have been subjected to veterinary controls that				
	are adapted to the intended use of the tissues. Information on the geographical origin of the animals shall be retained.				
	(b) Processing, preservation, testing and handling of tissues, cells and substances of				
	animal origin shall be carried out so as to provide optimal safety for patients, users				
	and, where applicable, other persons. In particular safety with regard to viruses and				
	other transmissible agents shall be addressed by implementation of validated				
	methods of elimination or viral inactivation in the course of the manufacturing				
	process.				
	(c) In the case of devices manufactured utilizing tissues of cells of animal origin as				
	narticular requirements as regards the requirements laid down in Council Directives				
	90/385/EEC and 93/42/EEC with respect to active implantable medical devices and				
	medical devices manufactured utilizing tissues of animal origin (OJ L 212, 9.8.2012,				
	p.3), the particular requirements laid down in that Regulation shall apply.				
10.3	For devices manufactured utilizing other non-viable biological substances the				
	following applies:				
	In the case of biological substances other than those referred to in Sections 10.1. and				
	10.2., the processing, preservation, testing and handling of those substances shall be				
	carried out so as to provide optimal safety for patients, users and, where applicable,				
	agents shall be addressed by implementation of validated methods of elimination or				
	inactivation in the course of the manufacturing process				
11	Interaction of devices with their environment				
11.1	If the device is intended for use in combination with other devices or equipment the				
	whole combination, including the connection system shall be safe and shall not impair				
	the specified performance of the devices. Any restrictions on use applying to such				
	combinations shall be indicated on the label and/or in the instructions for use.				
	Connections which the user has to handle, such as fluid, gas transfer or mechanical				
	coupling, shall be designed and constructed in such a way as to minimize all possible				

Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking Identity of the device and applicable configurations/variants covered by this checklist:



	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate	Method Reference	Reference to Supporting Controlled
			Conformity		Documents
	risks from incorrect connection.				
11.2	Devices shall be designed and manufactured in such a way as to remove or reduce				
	as far as possible and appropriate:				
	(a) the risk of injury to the patient, user or other persons in connection with their				
	physical and ergonomic features;				
	(b) the risk of use error due to the ergonomic features, human factors and the				
	environment in which the device is intended to be used;				
	(c) risks connected with reasonably foreseeable external influences or environmental				
	conditions, such as magnetic fields, external electrical and electromagnetic effects,				
	electrostatic discharge, radiation associated with				
	diagnostic or therapeutic procedures, pressure, numidity, temperature, variations in				
	pressure and acceleration or radio signal interferences;				
	(d) the risks associated with the use of the device when it comes into contact with				
	naterials, liquids, and substances, including gases, to which it is exposed during				
	(a) the risk appointed with the persible parative interaction between software and				
	(e) the fisk associated with the possible negative interaction between software and				
	(f) the risks of accidental ingress of substances into the device:				
	(i) the risks of accidental ingress of substances into the device,				
	(g) the fisks of recipiocal interference with other devices normally used in the				
	(h) risks arising where maintenance or calibration are not possible (as with implants)				
	from ageing of materials used or loss of accuracy of any measuring or control				
	mechanism				
11.3	Devices shall be designed and manufactured in such a way as to minimize the risks				
-	of fire or explosion during normal use and in single fault condition. Particular attention				
	shall be paid to devices whose intended purpose includes exposure to or use in				
	association with flammable substances or substances which could cause combustion.				
11.4	Devices shall be designed and manufactured in such a way that adjustment,				
	calibration, and maintenance, where such is necessary to achieve the performances				
	intended, can be done safely.				
11.5	Devices that are intended to be operated together with other devices or products shall				
	be designed and manufactured in such a way that the interoperability is reliable and				
-	safe.				
11.6	Any measurement, monitoring or display scale shall be designed in line with				
	ergonomic principles, taking account of the intended purpose of the device.				



	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate	Method Reference	Reference to Supporting Controlled
			Conformity		Documents
11.7	Devices shall be designed and manufactured in such a way as to facilitate the safe				
	disposal of the device and/or of any waste substances by the user, patient or other				
40	person.				
12	Devices with a diagnostic or measuring function				
12.1	Diagnostic devices and devices with a measuring function, shall be designed and manufactured in such a way as to provide sufficient accuracy, provision and stability				
	for their intended nurnese, based on appropriate scientific and technical methods				
	The limits of accuracy shall be indicated by the manufacturer				
12.2	The measurements made by devices with a measuring function and expressed in				
	legal units shall conform to the provisions of Council Directive 80/181/EEC (OJ L 39.				
	15.2.1980).				
13	Protection against radiation	1	1	1	
13.1	General				
	(a) Devices shall be designed and manufactured and packaged in such a way that				
	exposure of patients, users and other persons to any emitted radiation shall be				
	reduced as far as possible and appropriate, compatible with the intended purpose,				
	whilst not restricting the application of appropriate specified levels for therapeutic and				
	diagnostic purposes.				
	(b) The operating instructions for devices emitting radiation shall give detailed				
	information as to the nature of the emitted radiation, means of protecting the patient				
	and the user and on ways of avoiding misuse and of eliminating the risks inherent in				
42.0	Installation.				
13.2	(a) Where devices are designed to emit bezardous, or potentially bezardous, levels of				
	(a) where devices are designed to emit hazardous, or potentially hazardous, levels of visible and/or invisible radiation necessary for a specific medical nurpose the benefit				
	of which is considered to outweigh the risks inherent in the emission, it shall be				
	possible for the user to control the emissions. Such devices shall be designed and				
	manufactured to ensure reproducibility of relevant variable parameters within an				
	acceptable tolerance.				
	(b) Where devices are intended to emit potentially hazardous, visible and/or invisible				
	radiation, they shall be fitted, where possible, with visual displays and/or audible				
	warnings of such emissions.				
13.3	Unintended radiation				
	Devices shall be designed and manufactured in such a way that exposure of patients,				
	users and other persons to the emission of unintended, stray or scattered radiation is				

Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking Identity of the device and applicable configurations/variants covered by this checklist:



	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
	reduced as far as possible and appropriate.				
13.4	Ionizing radiation (a) Devices intended to emit ionizing radiation shall be designed and manufactured in such a way as to ensure that, where possible, the quantity, geometry and energy distribution (or quality) of radiation emitted can be varied and controlled taking into account the intended use.				
	designed and manufactured in such a way as to achieve appropriate image and/or output quality for the intended medical purpose whilst minimizing radiation exposure of the patient and user.				
	(c) Devices emitting ionizing radiation, intended for therapeutic radiology shall be designed and manufactured in such a way as to enable reliable monitoring and control of the delivered dose, the beam characteristics in terms of type of radiations, energy and, where appropriate, energy distribution.				
14	Software incorporated in devices and standalone software				
14.1	Devices that incorporate electronic programmable systems, including software, or standalone software that are devices in themselves, shall be designed to ensure repeatability, reliability and performance according to the intended use. In the event of a single fault condition, appropriate means shall be adopted to eliminate or reduce as far as possible and appropriate consequent risks.				
14.2	For devices that incorporate software or for standalone software that are devices in themselves, the software shall be developed and manufactured according to the state of the art taking into account the principles of development life cycle, risk management, verification and validation.				
14.3	Software referred to in this Section that are intended to be used in combination with mobile computing platforms shall be designed and manufactured taking into account the specific features of the mobile platform (e.g. size and contrast ratio of the screen) and the external factors related to their use (varying environment as regards to level of light or noise).				
15	Active devices and devices connected to them	1		[]	
15.1	For active devices, in the event of a single fault condition, appropriate means shall be adopted to eliminate or reduce as far as possible and appropriate consequent risks.				
15.2	Devices where the safety of the patients depends on an internal power supply shall be equipped with a means of determining the state of the power supply.				
15.3	Devices where the safety of the patients depends on an external power supply shall				

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Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking Identity of the device and applicable configurations/variants covered by this checklist:

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Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
include an alarm system to signal any power failure.				
Devices intended to monitor one or more clinical parameters of a patient shall be equipped with appropriate alarm systems to alert the user of situations which could lead to death or severe deterioration of the patient's state of health.				
Devices shall be designed and manufactured in such a way as to reduce as far as possible and appropriate the risks of creating electromagnetic interference which could impair the operation of this or other devices or equipment in the intended environment.				
Devices shall be designed and manufactured in such a way as to provide an adequate level of intrinsic immunity to electromagnetic disturbance to enable them to operate as intended.				
Devices shall be designed and manufactured in such a way as to avoid, as far as possible, the risk of accidental electric shocks to the patient, user or any other person, both during normal use of the device and in the event of a single fault condition in the device, provided the device is installed and maintained as indicated by the manufacturer.				
Protection against mechanical and thermal risks				
Devices shall be designed and manufactured in such a way as to protect the patient and user against mechanical risks connected with, for example, resistance to movement, instability and moving parts.				
Devices shall be designed and manufactured in such a way as to reduce to the lowest possible level the risks arising from vibration generated by the devices, taking account of technical progress and of the means available for limiting vibrations, particularly at source, unless the vibrations are part of the specified performance.				
Devices shall be designed and manufactured in such a way as to reduce to the lowest possible level the risks arising from the noise emitted, taking account of technical progress and of the means available to reduce noise, particularly at source, unless the noise emitted is part of the specified performance.				
Terminals and connectors to the electricity, gas or hydraulic and pneumatic energy supplies which the user or other person has to handle shall be designed and				

16.4 Te supp constructed in such a way as to minimize all possible risks. Errors likely to be made when fitting or refitting, or connecting or reconnecting, certain 16.5 parts before or during use which could be a source of risk must be made impossible by the design and construction of such parts or, failing this, by information given on the parts themselves and/or their housings.



	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
	The same information must be given on moving parts and/or their housings where the				
	direction of movement needs to be known in order to avoid a risk.				
16.6	Accessible parts of the devices (excluding the parts or areas intended to supply heat				
	or reach given temperatures) and their surroundings shall not attain potentially				
47	dangerous temperatures under normal conditions of use.				
17	Protection against the risks posed to the patient or user by supplied energy or su	Ibstances			
17.1	Devices for supplying the patient with energy or substances shall be designed and				
	constructed in such a way that the delivered amount can be set and maintained				
17.2	Devices shall be fitted with the means of preventing and/or indicating any				
17.2	inadequacies in the delivered amount which could pose a danger. Devices shall				
	incorporate suitable means to prevent as far as possible, the accidental release of				
	dangerous levels of energy or substances from an energy and/or substance source				
17.3	The function of the controls and indicators shall be clearly specified on the devices.				
	Where a device bears instructions required for its operation or indicates operating or				
	adjustment parameters by means of a visual system, such information shall be				
	understandable to the user and, as appropriate, the patient.				
18	Protection against the risks posed by medical devices intended by the manufactu	irer for use by lay	/ persons		
18.1	Devices for use by lay persons shall be designed and manufactured in such a way				
	that they perform appropriately for their intended purpose taking into account the				
	skills and the means available to lay persons and the influence resulting from				
	variation that can reasonably be anticipated in the lay person's technique and				
	environment. The information and instructions provided by the manufacturer shall be				
10.0	easy for the lay person to understand and apply.				
18.2	Devices for use by lay persons shall be designed and manufactured in such a way as				
	ensure that the device is easy to use by the intended user at all stages of the				
	procedure and				
	- reduce as far as possible the risk of error by the intended user in the handling of the				
	device and, if applicable, in the interpretation of the results.				
18.3	Devices for use by lay persons shall, where reasonably possible, include a procedure				
	by which the lay person				
	- can verify that, at the time of use, the device will perform as intended by the				
	manufacturer, and				
	<ul> <li>if applicable, is warned if the device has failed to provide a valid result.</li> </ul>				

Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking Identity of the device and applicable configurations/variants covered by this checklist:

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	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
	III. Requirements Regarding the Information Su	pplied with the D	evice		
19	Label and instructions for use				
19.1	General requirements regarding the information supplied by the manufacturer. Each device shall be accompanied by the information needed to identify the device and its manufacturer, and communicate safety and performance related information to the user, professional or lay, or other person, as appropriate. Such information may appear on the device itself, on the packaging or in the instructions for use, taking into account the following: (a) The medium, format, content, legibility, and location of the label and instructions for use shall be appropriate to the particular device, its intended purpose and the technical knowledge, experience, education or training of the intended user(s). In particular, instructions for use shall be written in terms readily understood by the intended user and, where appropriate, supplemented with drawings and diagrams.				
	Some devices may include separate information for the professional user and the lay				
	<ul><li>(b) The information required on the label shall be provided on the device itself. If this is not practicable or appropriate, some or all of the information may appear on the</li></ul>				
	Where multiple devices are supplied to a single user and/or location, a single copy of the instructions for use may be provided if so agreed by the purchaser who in any case may request further copies to be provided.				
	(c) For devices of class I and IIa, instructions for use are not needed or may be abbreviated if the device can be used safely and as intended by the manufacturer without any such instructions for use.				
	(d) Labels shall be provided in a human-readable format but may be supplemented by machine-readable forms, such as radio-frequency identification (RFID) or bar codes.				
	(e) Instructions for use may be provided to the user in non-paper format (e.g. electronic) to the extent and under the conditions set out in Commission Regulation (EU) No 207/2012 on electronic instructions for use of medical devices (OJ L 72, 10.3.2012, p.28).				
	(f) Residual risks which are required to be communicated to the user and/or other person shall be included as limitations, contraindications, precautions or warnings in the information supplied by the manufacturer.				
	(g) where appropriate, this information should take the form of internationally				



	Essential Requirements	Applicable to	Method Used to	Method	Reference to Supporting
		the Device?	Demonstrate Conformity	Reference	Controlled Documents
	recognized symbols. Any symbol or identification color used shall conform to the				
	harmonized standards or CTS. In areas for which no standards or CTS exist, the				
	symbols and colors shall be described in the documentation supplied with the device.				
19.2	Information on the label.				
	The label shall bear the following particulars:				
	(a) The name or trade name of the device.				
	(b) The details strictly necessary for a user to identify the device, the contents of the				
	packaging and, where it is not obvious for the user, the intended purpose of the				
	device.				
	(c) The name, registered trade name or registered trade mark of the manufacturer				
	and the address of his registered place of business at which he can be contacted and				
	(d) For imported devices, the name, registered trade name or registered trade mark				
	(d) For imported devices, the name, registered trade name or registered trade mark				
	of the authorized representative established within the Orifon and the authess of his				
	registered place of business at which he can be contacted and his location be				
	(a) Where applicable, an indication that the device contains or incorporates				
	a medicinal substance, including a human blood or plasma derivative, or				
	tissues or cells, or their derivatives, of human origin, or				
	tissues or cells, or their derivatives, of animal origin, or				
	Regulation (EU) No 722/2012.				
	(f) Where applicable, an indication that the device incorporates or consists of				
	nanomaterial unless the nanomaterial is encapsulated or bound in such a manner				
	that it cannot be released into the patient's or user's body when the device is used				
	within its intended purpose.				
	(g) The batch code/lot number or the serial number of the device preceded by the				
	word LOT or SERIAL NUMBER or an equivalent symbol, as appropriate.				
	(h) Where applicable, the unique device identification (UDI).				
	(I) An unambiguous indication of the date until when the device may be used safely,				
	expressed at least as the year and month, where this is relevant.				
	(j) where there is no indication of the date until when it may be used safely, the year				
	of manufacture. This year of manufacture may be included as part of the batch or				
	serial number, provided the date is clearly identifiable.				
	(k) An indication of any special storage and/or handling condition that applies.				
	(I) If the device is supplied sterile, an indication of its sterile state and the sterilization				

Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking Identity of the device and applicable configurations/variants covered by this checklist:



	Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
	method.				
	(m) Warnings or precautions to be taken that need to be brought to the immediate				
	attention of the user of the device as relevant, and to any other person where				
	appropriate. This information may be kept to a minimum in which case more detailed				
	information should appear in the instructions for use.				
	(n) If the device is intended for single use, an indication of that fact. A manufacturer's				
	indication of single use shall be consistent across the Union.				
	(o) If the device is a single use device that has been reprocessed, an indication of				
	that fact, the number of reprocessing cycles already performed, and any limitation as				
	regards the number of reprocessing cycles.				
	(p) If the device is custom made, an indication of that fact.				
40.0	(q) If the device is intended for clinical investigation only, an indication of that fact.				
19.3	Information in the instructions for use.				
	The instructions for use shall contain the following particulars: (a) The particulars referred to in points $10.2$ , a), b), c), f), k), l) and p)				
	(a) The particulars referred to in points 19.2. a), c), e), f), k), f) and f).				
	(b) The device's intended purpose including the intended user (e.g. professional of				
	(a) The performance of the device intended by the manufacturer				
	(d) Any residual risks, contraindications and any expected and foreseeable				
	(u) Any residual risks, contraindications and any expected and foreseeable				
	renard				
	(e) Specifications the user requires to use the device appropriately e.g. if the device				
	has a measuring function, the degree of accuracy claimed for it.				
	(f) Details of any preparatory treatment or handling of the device before it is ready for				
	use (e.g. sterilization, final assembly, calibration, etc.).				
	(g) Any requirements for special facilities, or special training, or particular				
	gualifications of the device user and/or other persons.				
	(h) The information needed to verify whether the device is properly installed and is				
	ready to perform safely and as intended by the manufacturer, together with, where				
	relevant:				
	- details of the nature, and frequency, of preventative and regular maintenance, and				
	of any preparatory cleaning or disinfection;				
	<ul> <li>identification of any consumable components and how to replace them;</li> </ul>				
	- information on any necessary calibration to ensure that the device operates				
	properly and safely during its intended lifetime;				

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Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
<ul> <li>methods of eliminating the risks encountered by persons involved in installing, calibrating or servicing devices.</li> </ul>				
(i) If the device is supplied sterile, instructions in the event of the sterile packaging being damaged before use.				
(j) If the device is supplied non-sterile with the intention that it is sterilized before use, the appropriate instructions for sterilization.				
(k) If the device is reusable, information on the appropriate processes to allow reuse, including cleaning, disinfection, decontamination, packaging, the maximum number of allowable reuses and, where appropriate, the validated method of re-sterilization. Information should be provided to identify when the device should no longer be reused, e.g. signs of material degradation or the maximum number of allowable reuses.				
(I) With the exception of devices referred to in Article 15b, if the device bears an indication that it is for single use, the evidence justifying that the device cannot be reprocessed safely referred to in Article 15 c (1), and which includes all information on characteristics and technical factors known to the manufacturer that could pose a risk if the device were to be re-used. If in accordance with point c) of Section 19.1 no instructions for use are needed, the information shall be made available to the user upon request.				
<ul> <li>(m) For devices intended for use together with other devices and/or general purpose equipment:</li> <li>– information to identify such devices or equipment, in order to obtain a safe combination, and/or</li> </ul>				
<ul> <li>information on any known restrictions to combinations of devices and equipment.</li> </ul>				
<ul> <li>(n) If the device emits hazardous, or potentially hazardous levels of radiation for medical purposes:</li> <li>– detailed information as to the nature, type and where appropriate, the intensity and distribution of the emitted radiation;</li> </ul>				
<ul> <li>the means of protecting the patient, user, or other person from unintended radiation during use of the device.</li> </ul>				
<ul> <li>(o) Information that allows the user and/or patient to be informed of any warnings, precautions, measures to be taken and limitations of use regarding the device. This information should cover, where appropriate:         <ul> <li>warnings, precautions and/or measures to be taken in the event of malfunction of the device or changes in its performance that may affect safety;</li> </ul> </li> </ul>				



Essential Requirements	Applicable to the Device?	Method Used to Demonstrate Conformity	Method Reference	Reference to Supporting Controlled Documents
<ul> <li>warnings, precautions and/or measures to be taken in regards to the exposure to reasonably foreseeable external influences or environmental conditions, such as magnetic fields, external electrical and electromagnetic effects, electrostatic discharge, radiation associated with diagnostic or therapeutic procedures, pressure, humidity, or temperature;</li> </ul>				
<ul> <li>warnings, precautions and/or measures to be taken in regards to the risks of interference posed by the reasonably foreseeable presence of the device during specific diagnostic investigations, evaluations, or therapeutic treatment or other procedures (e.g. electromagnetic interference emitted by the device affecting other equipment);</li> </ul>				
<ul> <li>if the device is intended to administer medicinal products, tissues or cells, or their derivatives, of human or animal origin or biological substances, any limitations or incompatibility in the choice of substances to be delivered;</li> </ul>				
<ul> <li>warnings, precautions and/or limitations related to the medicinal substance or biological material that is incorporated into the device as an integral part of the device;</li> </ul>				
<ul> <li>precautions related to materials incorporated into the device that are carcinogenic, mutagenic or toxic, or that have endocrine disrupting properties or that could result in sensitization or allergic reaction of the patient or user.</li> </ul>				
<ul> <li>(p) Warnings or precautions to be taken in order to facilitate the safe disposal of the device, its accessories and the consumables used with it, if any. This information should cover, where appropriate:</li> <li>– infection or microbial hazards (e.g. explants, needles or surgical equipment contaminated with potentially infectious substances of human origin);</li> </ul>				
<ul> <li>physical hazards (e.g. from sharps).</li> <li>(q) For devices intended for use by lay persons, the circumstances when the user</li> </ul>				
<ul> <li>should consult with a healthcare professional.</li> <li>(r) For devices listed in Annex XV for which the manufacturer does not claim a medical purpose, information regarding the absence of a clinical benefit and the risks related to the use of the device.</li> </ul>				
(s) Date of issue of the instructions for use or, if they have been revised, date of issue and identifier of the latest revision of the instructions for use.				
(t) A notice to the user and/or patient that any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State where the user and/or patient is established.				

**Essential Requirements Checklist** Annex I of Proposed EU Regulations & Compromise Amendment for Medical Device CE Marking

Identity of the device and applicable configurations/variants covered by this checklist:



References:

- 1. Proposed EU Medical Device Regulations. European Commission. September 26, 2012.
- 2. Compromise and Consolidated Amendments 1-30, Draft Report. Dagmar Roth-Behrendt. September 23, 2013. (Summary of Changes: 19.2(o) deleted, 19.3(k) edited, 19.3(l) edited.)