






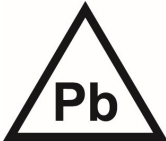




















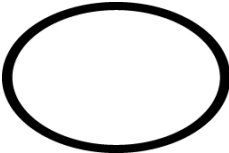
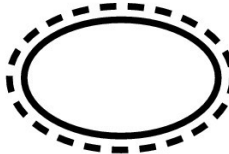
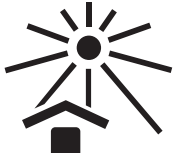




























| REFERENCE-NUMMER   | SYMBOL  | FORKLARENDE TEKST/FORKLARENDE TEKST  |
|--|---|--|
| <b>INGEN STANDARDREFERENCE/IKKE ET UNIVERSALT ANERKENDT ELLER STANDARDISERET SYMBOL</b>  |   |  |
|  |    | Ansvarlig person Storbritannien  |
|  |    | Mængde   |
|  |    | Udstyret indeholder ikke og anvender ikke latex/naturgummi                           |
|  |    | Kommentarer fra kunder   |
|  |   | Læs venligst de vigtige sikkerhedsrelaterede oplysninger i de medfølgende dokumenter |
|  |  | Mærket som type CF udstyr med hensyn til patientbeskyttelse                          |
|  |  | Indeholder CMR-stoffet kobolt  |
|  |  | Indeholder CMR-stoffet, bly  |
|  |  | Indeholder nikkel  |
|  |  | Indeholder dioctyltin bis(2-ethylhexylmercaptoacetat)                                |
| <b>ISO 15223-1 - MEDICINSKE UDSTYR - SYMBOLER, DER SKAL ANVENDES PÅ ETIKETTER, MÆRKNING OG INFORMATION, DER SKAL LEVERES MED MEDICINSKE UDSTYR</b> |   |  |
| 5.1.1  |  | Producent  |

| REFERENCE-NUMMER | SYMBOL  | FORKLARENDE TEKST/FORKLARENDE TEKST                                       |
|------------------|---|---|
| 5.1.2            |    | Autoriseret repræsentant i Det Europæiske Fællesskab/Den Europæiske Union |
| 5.1.3            |    | Produktionsdato   |
| 5.1.4            |    | Anvendes inden  |
| 5.1.5            |    | Batch-kode  |
| 5.1.6            |    | Katalognummer   |
| 5.1.7            |   | Serienummer   |
| 5.1.8            |  | Importør  |
| 5.1.9            |  | Distributør   |
| 5.1.11           |  | Fremstillingsland – Mexico  |
| 5.1.11           |  | Fremstillingsland – USA   |
| 5.2.3            |  | Steriliseret med ethylenoxid  |
| 5.2.3/5.2.11     |  | Steriliseret med ethylenoxid/System med en enkelt steril barriere         |

| REFERENCE-NUMMER | SYMBOL  | FORKLARENDE TEKST/FORKLARENDE TEKST  |
|------------------|---|--|
| 5.2.3/5.2.14     |    | Steriliseret med ethylenoxid/Enkeltvist system med steril barriere og beskyttende ydre emballering |
| 5.2.4            |    | Steriliseret ved brug af bestråling  |
| 5.2.4/5.2.11     |    | Steriliseret ved brug af bestråling/System med en enkelt steril barriere                           |
| 5.2.6            |    | Må ikke resteriliseres   |
| 5.2.8            |    | Må ikke anvendes, hvis indpakningen er beskadiget og se brugsanvisningen                           |
| 5.2.11           |  | System med en enkelt steril barriere   |
| 5.2.14           |  | Enkeltvist system med steril barriere og beskyttende ydre emballering                              |
| 5.3.2            |  | Udsæt ikke for sollys  |
| 5.3.4            |  | Opbevares tørt   |
| 5.3.6            |  | Øvre temperaturgrænse  |
| 5.3.7            |  | Temperaturgrænse   |

| REFERENCE-NUMMER | SYMBOL   | FORKLARENDE TEKST/FORKLARENDE TEKST                          |
|------------------|--|--|
| 5.4.2            |                     | Må ikke genbruges  |
| 5.4.3            | <br>www.medcomp.net | Se brugsanvisningen eller se brugsanvisningen for elektronik |
| 5.4.4            |                     | Forsigtig  |
| 5.4.7            |                     | Indeholder et lægemiddel                                     |
| 5.4.8            |                    | Indeholder biologisk materiale fra dyr                       |
| 5.4.10           |                   | Indeholder skadelige stoffer                                 |
| 5.6.3            |                   | Ikke-pyrogen   |
| 5.7.3            |                   | Patientidentifikation  |
| 5.7.4            |                   | Hjemmeside med oplysninger til patienter                     |
| 5.7.5            |                   | Sundhedscenter eller læge                                    |
| 5.7.6            |                   | Dato   |

| REFERENCE-NUMMER   | SYMBOL  | FORKLARENDE TEKST/FORKLARENDE TEKST   |
|--|---|---|
| 5.7.7  |    | Medicinsk udstyr  |
| 5.7.10   |    | Unik enhedsidentifikationskode  |
| <b>BS EN 15986 - SYMBOL TIL ANVENDELSE TIL MÆRKNING AF MEDICINISKE UDSTYR, KRAV TIL MÆRKNING AF MEDICINSK UDSTYR, DER INDEHOLDER PHTALAT</b> |   |   |
| A.2  |    | Indeholder ftalat: DEHP   |
| <b>FDA-VEJLEDNING ANVENDELSE AF SYMBOLER I MÆRKNING (CFR 801)</b>  |   |   |
|  | <b>Rx Only</b>  | Advarsel: Føderal lovgivning begrænser denne enhed til salg af eller efter ordre fra en autoriseret læge. |
| <b>ASTM F2503 - STANDARDPRAKSIS FOR MÆRKNING AF MEDICINSK UDSTYR OG ANDRE ENHEDER FOR SIKKERHED I MILJØ MED MAGNETISK RESONANS</b>           |   |   |
|  |   | MR-sikker   |
|  |  | Betinget MR-sikker  |
|  |  | MR-usikker  |
|  |  | Betinget MR-sikker – 3,0 Tesla  |
|  |  | Betinget MR-sikker – 1,5, 3,0 Tesla   |
| <b>ISO 7000 - GRAFISKE SYMBOLER TIL BRUG PÅ UDSTYRSREGISTREREDE SYMBOLER</b>   |   |   |
| 3079   |  | Åbnes her   |

| REFERENCE-NUMMER   | SYMBOL  | FORKLARENDE TEKST/FORKLARENDE TEKST |
|--|---|-------------------------------------|
| <b>MDD 93/42/EEC, MDR 2017/745, FORORDNING (EF) 765/2008</b> |   |                                     |
|  |  | CE-overensstemmelsesmærkning        |
| <b>LOV AF 2021 FOR MEDICINER OG MEDICINISKE UDSTYR</b>       |   |                                     |
|  |  | UK overensstemmelse vurderet        |
| <b>SWISSMEDIC-REGULATIVER</b>                                |   |                                     |
|  |  | Autoriseret repræsentant i Schweiz  |