














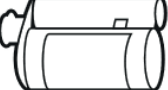

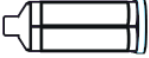

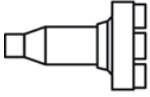






















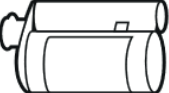
Explanation of symbols used by Sultan Healthcare	English	2
Explication des symboles utilisés par Sultan Healthcare	Français	5


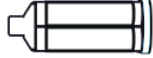

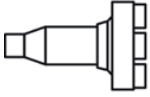





	<p>Medical device</p> <p>Indicates the item is a medical device. ISO 15223-1.</p>		<p>Caution</p> <p>Indicates that caution is necessary when operating the device or control close to where the symbol is placed, or that the current situation needs operator awareness or operator action in order to avoid undesirable consequences. ISO 7000-0434A, ISO 15223-1.</p>
	<p>Manufacturer</p> <p>Indicates the medical device manufacturer. ISO 7000-3082, ISO 15223-1.</p>		<p>Keep away from sunlight</p> <p>Indicates a medical device that needs protection from light sources. ISO 7000-0615, ISO 15223-1.</p>
	<p>Rx only</p> <p>Indicates potential for harmful effect and use is not safe without supervision of a practitioner. USA Code of Federal Regulations 21 CFR Part 801 § 801.109 (b) (1)</p>		<p>Do not use if package is damaged and consult instructions for use</p> <p>Indicates that a medical device that should not be used if the package has been damaged or opened and that the user should consult the instructions for use for additional information. ISO 7000-2606, ISO 15223-1.</p>
	<p>Use-by date</p> <p>Indicates the date after which the medical device is not to be used- ISO 7000-2607, ISO 15223-1.</p>		<p>Keep dry</p> <p>Indicates a medical device that needs to be protected from moisture. ISO 7000-0626, ISO 15223-1.</p>

	<p>Batch code</p> <p>Indicates the manufacturer's batch code so that the batch or lot can be identified. ISO 7000-2492, ISO 15223-1.</p>		<p>Temperature limit</p> <p>Indicates the temperature limits to which the medical device can be safely exposed. ISO 7000-0632, ISO 15223-1.</p>
	<p>Catalogue number</p> <p>Indicates the manufacturer's catalogue number (SKU) so that the medical device can be identified. ISO 7000-2493, ISO 15223-1.</p>		<p>Contains nano materials</p> <p>Indicates a medical device that contains nano materials. ISO 7000-3703, ISO 15223-1.</p>
	<p>Do not reuse</p> <p>Indicates a medical device that is intended for one single use only or for use on a single patient during a single procedure. ISO 7000-1051, ISO 15223-1.</p>		<p>Intraoral tip</p>
	<p>Date of manufacture</p> <p>Indicates the date when the medical device was manufactured ISO 7000-2497; ISO 15223-1.</p>		<p>380ml Cartridge</p>

	<p>Dispenser</p>		<p>50ml Cartridge</p>
	<p>Bayonet Ring</p>		<p>380ml Mixing tip</p>
	<p>50ml Mixing tip</p>		<p>Berry aroma Indicates the berry aroma</p>
	<p>Standard set clock Followed by the specific minutes, it indicates the set timings.</p>		<p>Mint aroma Indicates the mint aroma</p>
	<p>Rapid set clock or Fast/Super Fast set clock (only for Bite) Followed by the specific minutes, it indicates the set timings.</p>		

	<p>Dispositif médical</p> <p>Indique que l'article est un dispositif médical. ISO 15223-1.</p>		<p>Attention</p> <p>Indique que la prudence est nécessaire lors de l'utilisation du dispositif ou de la commande près de l'emplacement où se trouve le symbole, ou que la situation actuelle exige une sensibilisation de l'opérateur afin d'éviter des conséquences indésirables. ISO 7000-0434A, ISO 15223-1.</p>
	<p>Fabricant</p> <p>Indique le fabricant du dispositif médical. ISO 7000-3082, ISO 15223-1.</p>		<p>Conserver à l'abri de la lumière du soleil</p> <p>Indique un dispositif médical qui a besoin d'être protégé des sources de lumière. ISO 7000-0615, ISO 15223-1.</p>
	<p>Sur prescription uniquement</p> <p>Indique qu'il existe un potentiel d'effet nocif et que l'utilisation n'est pas sécuritaire sans la supervision d'un praticien. Code of Federal Regulations des États-Unis 21 CFR Part 801 § 801.109 (b) (1)</p>		<p>Ne pas utiliser si l'emballage est endommagé et consulter le mode d'emploi</p> <p>Indique qu'un dispositif médical ne devrait pas être utilisé si l'emballage a été endommagé ou ouvert et que l'utilisateur devrait consulter le mode d'emploi pour plus de renseignements. ISO 7000-2606, ISO 15223-1.</p>
	<p>Date limite d'utilisation</p> <p>Indique la date après laquelle le dispositif médical ne doit plus être utilisé. ISO 7000-2607, ISO 15223-1.</p>		<p>Conserver au sec</p> <p>Indique un dispositif médical qui a besoin d'être protégé de l'humidité. ISO 7000-0626, ISO 15223-1</p>

	<p>Code du lot</p> <p>Indique le code du lot du fabricant permettant d'identifier le lot. ISO 7000-2492, ISO 15223-1.</p>		<p>Limite de température</p> <p>Indique les limites de température auxquelles le dispositif médical peut être exposé de manière sécuritaire. ISO 7000-0632, ISO 15223-1.</p>
	<p>Référence catalogue</p> <p>Indique la référence catalogue du fabricant (numéro de référence) permettant d'identifier le dispositif médical. ISO 7000-2493, ISO 15223-1</p>		<p>Contient des nanomatériaux</p> <p>Indique un dispositif médical qui contient des nanomatériaux. ISO 7000-3703, ISO 15223-1.</p>
	<p>Ne pas réutiliser</p> <p>Indique un dispositif médical prévu pour un usage unique ou pour une utilisation chez un seul patient au cours d'une seule procédure. ISO 7000-1051, ISO 15223-1.</p>		<p>Embout intra-oral</p>
	<p>Date de fabrication</p> <p>Indique la date à laquelle le dispositif médical a été fabriqué ISO 7000-2497; ISO 15223-1.</p>		<p>Cartouche de 380 ml</p>

	<p>Distributeur</p>		<p>Cartouche de 50 ml</p>
	<p>Anneau de baïonnette</p>		<p>Embout mélangeur de 380 ml</p>
	<p>Embout mélangeur de 50 ml</p>		<p>Arôme de baies Indique que le produit est aromatisé aux baies</p>
	<p>Prise standard Suivi d'un nombre de minutes, indique le temps de prise du produit.</p>		<p>Arôme de menthe Indique que le produit est aromatisé à la menthe</p>
	<p>Prise rapide ou Prise rapide/super rapide (seulement pour occlusion) Suivi d'un nombre de minutes, indique le temps de prise du produit.</p>		