

Tristel™

TRISTEL DUO OPH

Rapid and reliable
high-level disinfection
for ophthalmic devices.

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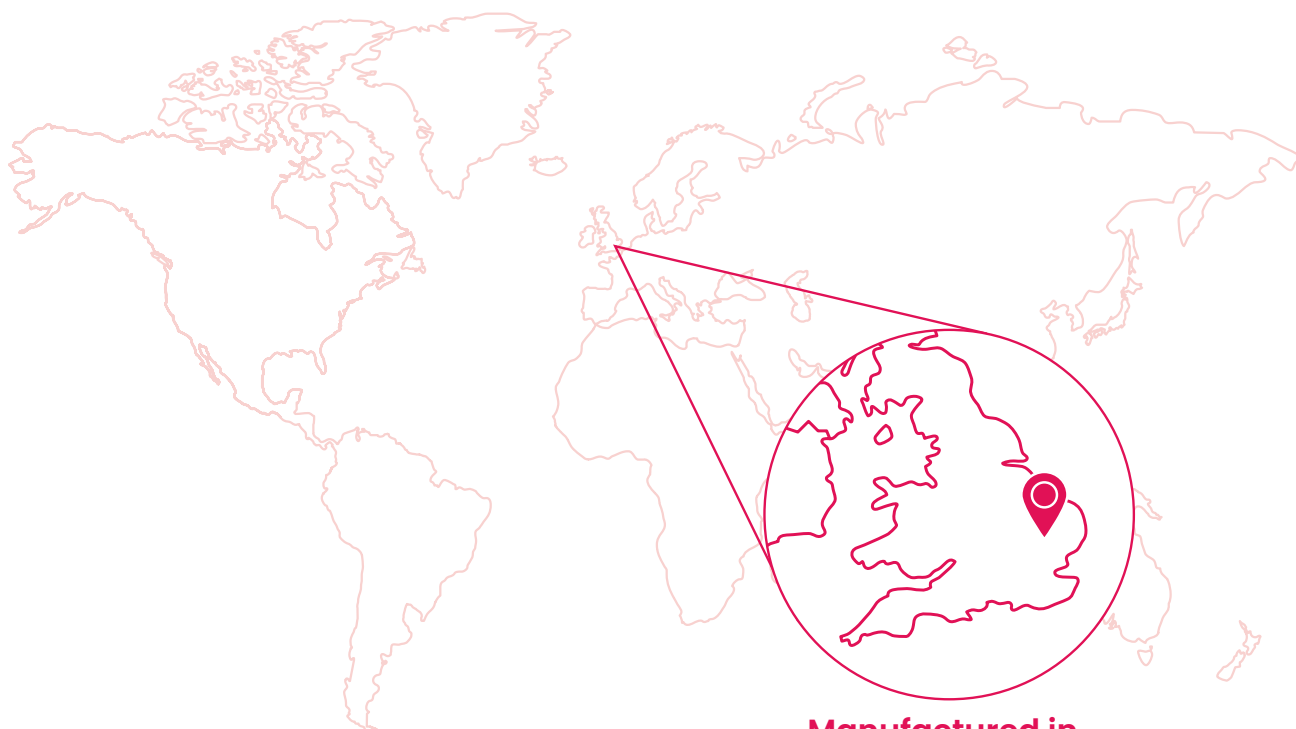
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> **Tristel chlorine dioxide has been used in an estimated 100 million+ decontamination procedures**

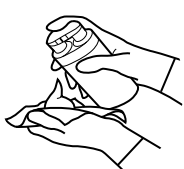
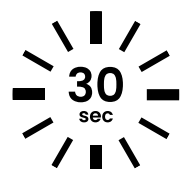
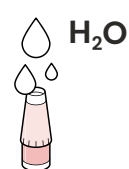
Tristel DUO OPH provides high-level disinfection of ophthalmic medical devices, such as diagnostic contact lenses, tonometer prisms, ophthalmic ultrasound probes and handheld pachymeters. Proven to be sporicidal, mycobactericidal, virucidal, fungicidal, yeasticidal and bactericidal in only 30 seconds, Tristel DUO OPH offers quick, effective and mobile protection against even the most difficult-to-eliminate microorganisms.



**Manufactured in
Cambridgeshire, UK**

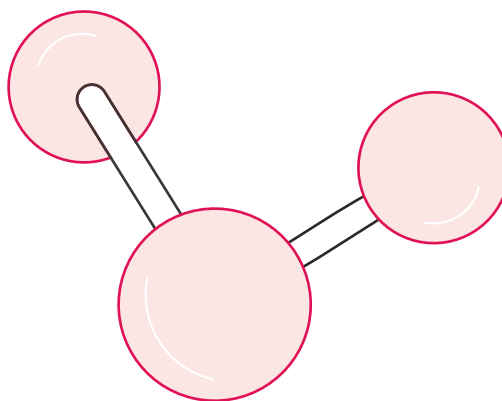
How does it work?

Tristel DUO OPH can be used with Tristel DUO WIPES and 3T to provide you with a full ophthalmology decontamination process with traceability.

HIGH-LEVEL DISINFECT				TRACE
				
Dispense Tristel DUO OPH onto a Tristel DUO WIPE.	Wipe your device.	Observe the contact time.	Rinse the device.	Track your decontamination cycles with 3T.

 Refer to user guide for full instructions.

WE HAVE CHEMISTRY



Tristel Chlorine Dioxide

Tristel's proprietary chlorine dioxide (ClO_2) chemistry is trusted globally in healthcare settings for its fast-acting, easy-to-use and effective disinfection across diverse medical fields.

ClO_2 kills pathogens through electron exchange, stealing electrons from the microorganism's structures. Due to this reaction mechanism, microorganisms cannot develop resistance.

Tristel's chemistry is designed to work with innovative delivery systems to facilitate simple, but effective point of use disinfection, ensuring exceptional efficacy. Tristel's proprietary chlorine dioxide chemistry has a broad spectrum of biocidal efficacy and is proven effective against bacteria, bacterial spores, mycobacteria, enveloped and non-enveloped viruses, fungi and yeast.



Broad Spectrum



Fast Acting



Ease of Use



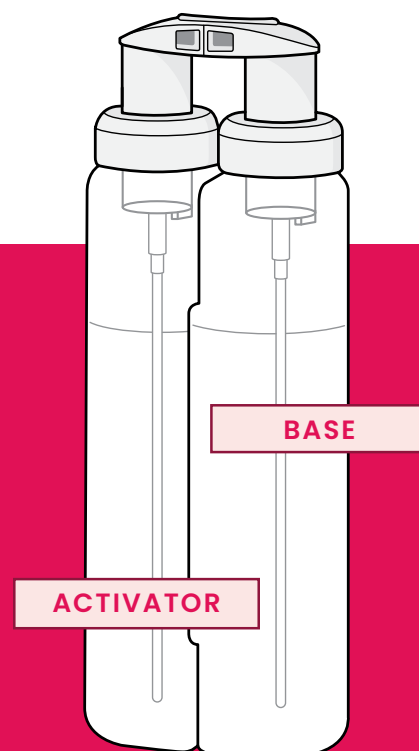
Cleaning Properties



Prevents the Spread of AMR Pathogens

> Tristel DUO OPH is free from alcohol and Quaternary Ammonium Compounds (QAC)

Tristel DUO OPH is simple: It has two separate compartments that contain 125ml of Tristel Base Solution (citric acid) and 125ml of Tristel Activator Solution (sodium chlorite). When the pump is pressed, the two solutions combine and chlorine dioxide chemistry is generated as a foam, ready to disinfect.



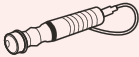
WHY HIGH-LEVEL DISINFECT?

The Spaulding Classification

Decontamination of medical devices is critical when it comes to preventing hospital acquired infections (HAIs), but why should you high-level disinfect ophthalmic devices?

The Spaulding Classification determines the appropriate level of disinfection (critical, semi-critical and non-critical) for medical devices, depending on the degree of risk of infection when used.¹

All ophthalmic devices which come into contact with non-intact skin or mucous membrane should be high-level disinfected.

CATEGORY	DEVICE APPLICATION		REQUIRED LEVEL OF DISINFECTION
CRITICAL	Contact with the bloodstream or sterile tissues.	 Surgical instruments, e.g. scalpels, tweezers, scissors, and clamps.	Sterilisation Eliminates all forms of microbial life.
SEMI-CRITICAL	Contact with mucous membranes or non-intact skin.	 Contact lenses, tonometer prisms, ophthalmic ultrasound probes, handheld pachymeters.	High-Level Disinfection Destroys all vegetative microorganisms, mycobacteria, enveloped and non-enveloped viruses, fungal spores and some bacterial spores.
NON-CRITICAL	Contact with intact skin.	 Ultrasound Probes which do not come into contact with non-intact skin or mucous membrane.	Intermediate-Level Disinfection Destroys mycobacteria, most viruses, most fungi and bacteria.
		 Stethoscopes and blood pressure cuffs.	Low-Level Disinfection Destroys most bacteria, some viruses and some fungi.

Please note Tristel DUO OPH is a high-level disinfectant and is only suitable for the disinfection of semi-critical and non-critical devices.

EXCEPTIONAL EFFICACY

Effective in 30 seconds



Tristel DUO OPH is a high-level disinfectant, proven effective against a wide range of hard-to-kill microorganisms in **only 30 seconds**. All Tristel products are extensively tested according to relevant European tests such as those specified within the EN 14885.

STANDARD	ORGANISM TYPE	ORGANISM	TEST CONDITIONS
EN 17846	Bacterial Spores	<i>Clostridioides difficile</i>	Clean
			Dirty
EN 17126	Bacterial Spores	<i>Bacillus subtilis</i>	Clean
			Dirty
		<i>Bacillus cereus</i>	Clean
			Dirty
		<i>Clostridioides difficile</i>	Clean
			Dirty
EN 14348	Mycobacteria	<i>Mycobacterium terrae</i>	Clean
			Dirty
		<i>Mycobacterium avium</i>	Clean
			Dirty
EN 14476	Viruses	Poliovirus	Clean
			Dirty
		Adenovirus	Clean
			Dirty
		Murine Norovirus	Clean
			Dirty
EN 13624	Fungi	<i>Aspergillus brasiliensis</i>	Clean
			Dirty
	Yeasts	<i>Candida albicans</i>	Clean
			Dirty

According to the acceptance criteria of the European standard: Bacterial spores, mycobacteria, fungi, yeast and viruses: $\geq 4 \log_{10}$ reduction. Bacteria: $\geq 5 \log_{10}$ reduction. Additional requirement for 4-field tests: F2-F4 $< 50 \text{ cfu/cm}^2$

EXCEPTIONAL EFFICACY, CONTINUED

STANDARD	ORGANISM TYPE	ORGANISM	TEST CONDITIONS
EN 16615	Yeasts	<i>Candida albicans</i>	Clean
			Dirty
	Bacteria	<i>Staphylococcus aureus</i>	Clean
			Dirty
		<i>Pseudomonas aeruginosa</i>	Clean
			Dirty
		<i>Enterococcus hirae</i>	Clean
			Dirty
EN 13727	Bacteria	<i>Staphylococcus aureus</i>	Clean
			Dirty
		<i>Pseudomonas aeruginosa</i>	Clean
			Dirty
		<i>Enterococcus hirae</i>	Clean
			Dirty

According to the acceptance criteria of the European standard: Bacterial spores, mycobacteria, fungi, yeast and viruses: $\geq 4 \log_{10}$ reduction.
 Bacteria: $\geq 5 \log_{10}$ reduction. Additional requirement for 4-field tests: F2-F4 $< 50 \text{ cfu/cm}^2$



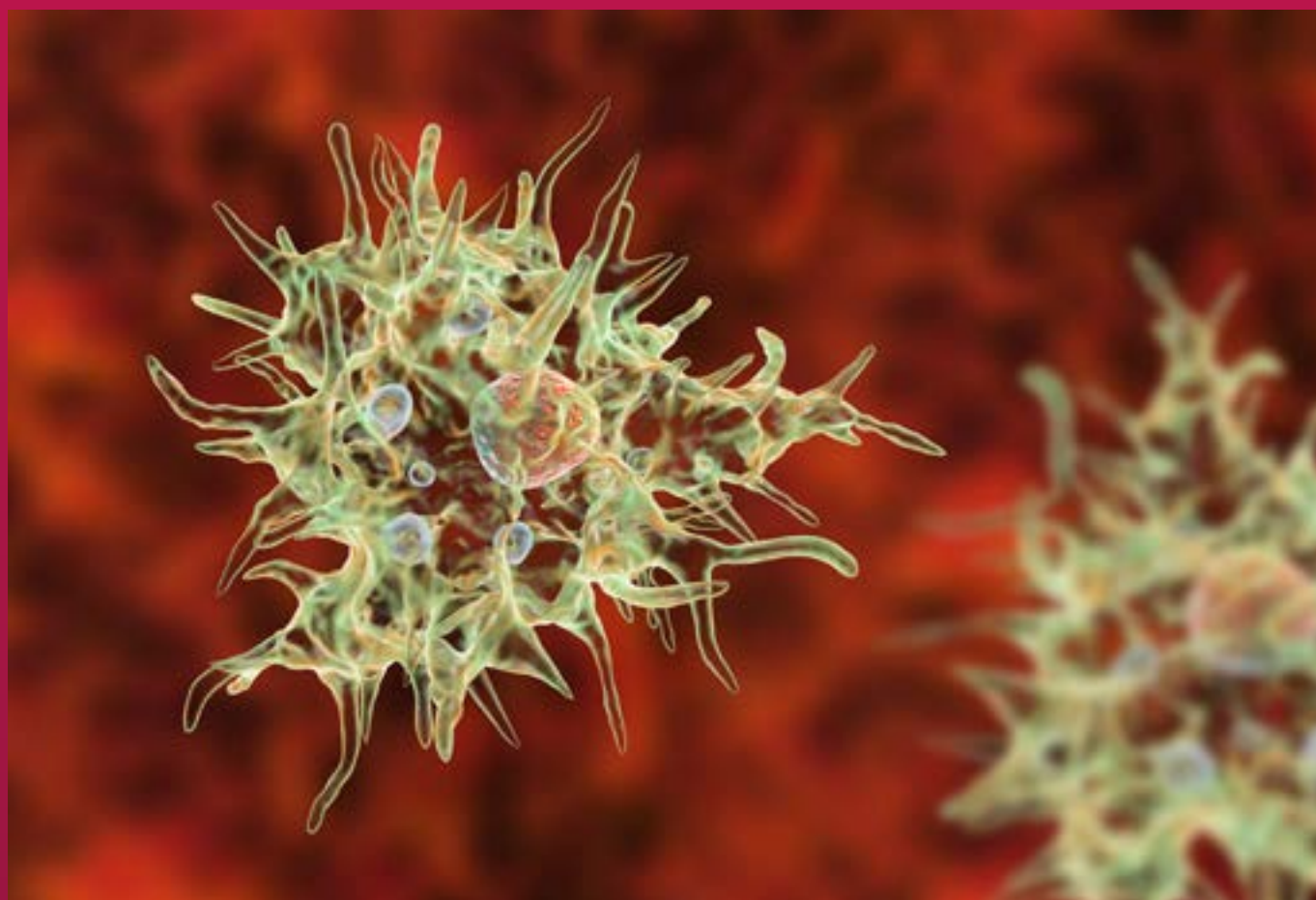
PROTECT YOUR PATIENTS

Against Priority Pathogens

Ophthalmic devices are often exposed to potentially harmful microorganisms because they are used near or in direct contact with the eye's surface. This exposure elevates the risk of transmitting dangerous pathogens that may lead to serious infections such as conjunctivitis, keratitis, and endophthalmitis.

Acanthamoeba castellanii cysts

Acanthamoeba is a free-living amoeba frequently found in soil, water, and occasionally in contact lens solutions. Its cyst form is notably resistant to environmental stressors. The primary concern with this organism is its association with *Acanthamoeba* keratitis, a serious infection that occurs when amoeba infects the cornea. If left untreated, this condition can lead to significant eye damage and permanent vision loss.



Tristel DUO OPH has been tested for its effectiveness to eliminate *Acanthamoeba castellanii* cysts at an ISO 17025 accredited laboratory, where Tristel DUO OPH demonstrated a $>3 \log_{10}$ reduction and a complete inactivation of *Acanthamoeba castellanii* cysts.

PROTECT YOUR PATIENTS

Against Priority Pathogens

Tristel DUO OPH also effectively eliminates:



Adenovirus

The primary cause of viral conjunctivitis, accounting for approximately 65–95% of all cases.² Highly contagious, it can be transmitted through direct contact, contaminated surfaces and ophthalmic instruments used during eye examinations.



Neisseria gonorrhoeae

N. gonorrhoeae can cause Gonococcal conjunctivitis (GC), a serious condition that can lead to complications like blindness or systemic infection. Approximately 10% of neonates exposed to fluids contaminated with *N. gonorrhoeae* during delivery may go on to develop GC.⁴

Candida albicans

Candida species are among the most frequent microorganisms associated with fungal infections (candidiasis) such as keratitis and endophthalmitis, and candidemia. One study found that the incidence of ocular candidiasis in patients with candidemia ranged from 2–26%.⁷



Fusarium solani

Fusarium keratitis is a severe ocular infection, caused by the organism *Fusarium solani*, it is a common cause of monocular blindness. The annual prevalence of fungal keratitis is estimated to be over 1 million cases globally. Among these cases, *Fusarium* species is the most frequently isolated cause.^{7,8}



Pseudomonas aeruginosa

Globally, an estimated 10–15% of nosocomial infections are due to *P. aeruginosa*. In addition, it is the most identified causative organism in contact lens-related keratitis.⁵

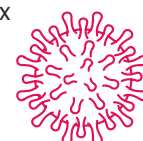


Staphylococcus aureus

S. aureus is a common cause of ocular infections such as conjunctivitis, keratitis, and endophthalmitis. Approximately 35% of the general public and 50–66% of hospital workers become colonised with *S. aureus*.⁶

Herpes simplex virus (HSV)

Can cause a range of diseases, including Herpes Simplex Keratitis (HSK), or ocular herpes, an infection that can lead to serious eye complications. HSV is estimated to cause 1.5 million cases annually, with 40,000 new cases of severe monocular visual impairment or blindness each year.³



PROTECT YOUR PATIENTS

Against Priority Pathogens – AMR

Antimicrobial resistance (AMR) is a critical global healthcare challenge, as microorganisms continue to evolve, rendering treatments for common infections less effective. This leads to increased healthcare costs, prolonged patient recovery times, and higher mortality rates.

Based on estimates across 204 countries and territories, new forecasts from the Global Research on Antimicrobial Resistance (GRAM) Project suggest that bacterial antimicrobial resistance (AMR) will cause **39 million deaths between 2025 and 2050 – which equates to three deaths every minute.**⁹

Tristel DUO OPH has been specifically tested against pathogens with known antibiotic resistance mechanisms, helping to prevent the spread of antimicrobial resistant organisms.

ClO_2 kills pathogens through electron exchange, stealing electrons from the microorganism's structures. **Due to this reaction mechanism microorganisms cannot develop resistance.**

Tristel DUO OPH is effective in 30 seconds against:



***Clostridioides
difficile***



**Methicillin-resistant
*Staphylococcus
aureus* (MRSA)**



**Carbapenem-resistant
Enterobacteriaceae (CRE)
*Klebsiella pneumoniae***



**Multidrug-resistant
*Acinetobacter
baumannii* (MDRAB)**



**Extended Spectrum
Beta-Lactamase
Klebsiella pneumoniae (ESBL)**



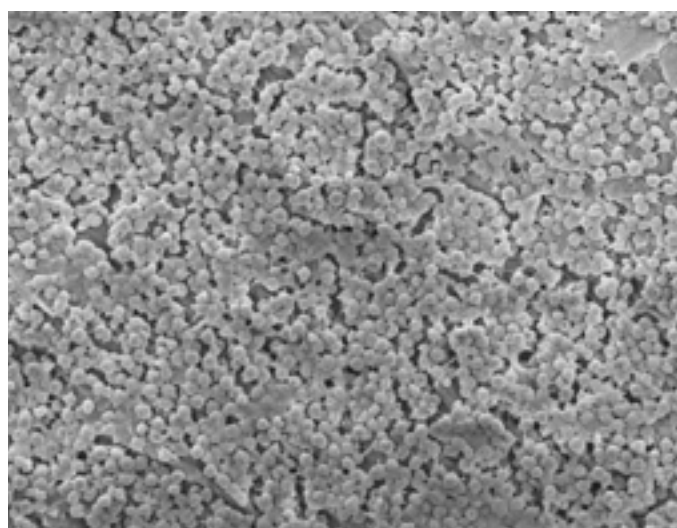
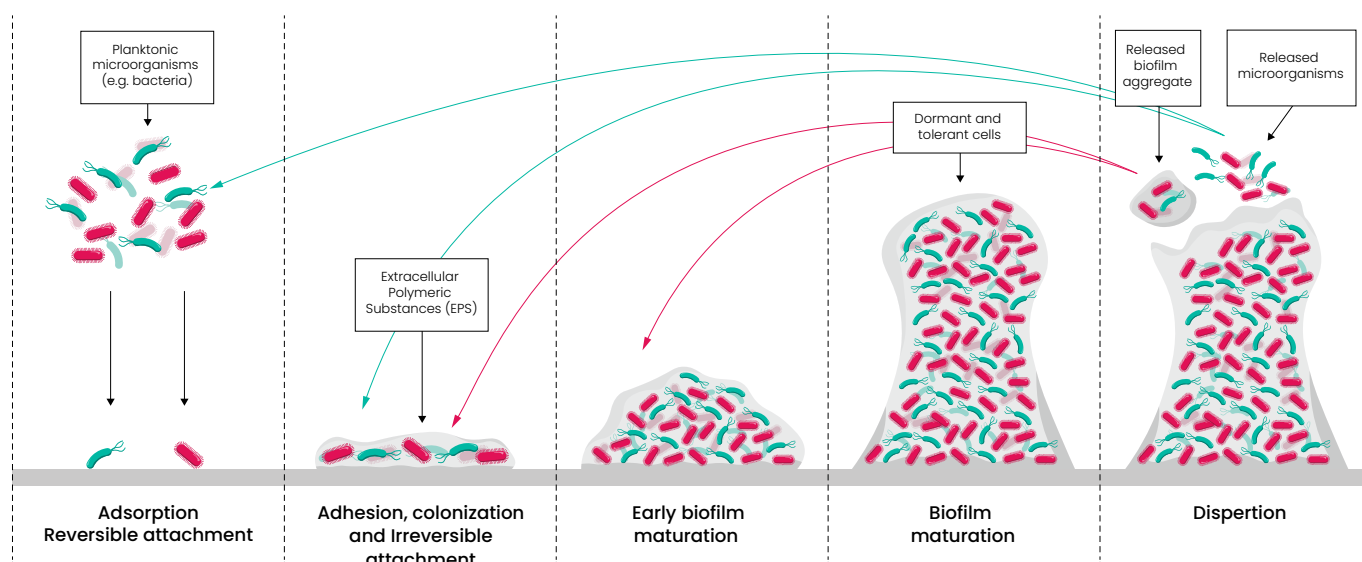
**Vancomycin-resistant
Enterococci (VRE)
*Enterococcus faecium***

PROTECT YOUR PATIENTS

Against Priority Pathogens – Biofilms

Biofilms are a significant issue in hospitals, they can provide a protective environment for microorganisms, allowing them to survive in harsh conditions, including exposure to disinfectants and antibiotics. These complex communities of microorganisms adhere to surfaces such as medical devices and general surfaces, making the microorganisms particularly difficult to eliminate.

Bacteria living in a biofilm exhibit a 10 to 1,000-fold increase in resistance to antibiotics compared to their planktonic counterparts.¹⁰



Biofilms can lead to persistent infections, increased resistance to treatments and a heightened risk of cross-contamination. Their presence on medical equipment, environmental surfaces and within environments such as water systems can also contribute to hospital-acquired infections (HAIs), posing a serious risk to patient safety.

It's estimated that around 65–80% of Hospital Acquired Infections are linked to biofilms.^{11,12}

Tristel DUO OPH has been specifically tested for its efficacy against both wet and dry biofilms, ensuring your product is effective in these environments.

COMPATIBILITY

With Major Manufacturers

Tristel DUO OPH has been tested and proven to be compatible with the instruments of major manufacturers, including:

- DGH Technologies
- Ellex
- Haag-Streit
- Keeler Accutome
- Laboratoires Thea
- Natus Medical
- Neolight / Phoenix Technology Group
- Nidek
- Ocular Instruments
- Quantel Medical
- Reichert Technologies
- Takagi
- Tomey
- Volk





DIGITAL TRACEABILITY & TRAINING

Say goodbye to paper-based traceability



Complete

Cloud-based traceability and training platform



Compatible

With
Tristel DUO OPH



Compliant

Be compliant with guidelines by recording your decontamination procedures with 3T

Tristel DUO OPH is fully compatible with 3T, Tristel's cloud-based compliance platform, created to guide you through the decontamination process and provide you with greater visibility on your infection control procedures.

Recording your Tristel DUO OPH decontamination processes through 3T ensures your procedures are fully traceable.

Other features of 3T include:

- Product training and certifications
- Secure administration portal
- User-friendly and accessible dashboards
- Localisation and scan features



HOW TO ORDER

Tristel DUO OPH

High-level disinfectant foam



Tristel DUO WIPES

Soft, durable, low-linting wipes

Ordering information:

Tristel DUO OPH
Product Code: TSL022701

Tristel DUO WIPES
Product Code: TSL031601

Additional products:

Tristel Pre-Clean Wipes
Product Code: TSL030401

Tristel CLEAN
Product Code: TSL023301

Tristel Rinse Wipes
Product Code: TSL030301

Tristel DUO OPH is classified as Class IIa Medical Device according to UK MDR and EU MDR.
 Tristel DUO WIPES are classified as a Class I Medical Device according to UK MDR and EU MDR.

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