



IV THERAPY
Closed Needleless Connectors



vadsite[®]
The Transparent, Split
Septum, Needleless
Connector



Value Life

Why choose **vadsite**?

vadsite is a closed, needleless connector for use with all I.V. equipment for example: vascular access devices, syringes, giving sets, stopcocks, extension lines, catheters and cannulae.

When deciding which needleless connector to choose for your hospital, it is important to ensure that your choice meets the current 'global standards' for these devices. The global opinion leaders make a number of recommendations in terms of the essential features you should demand when choosing a needleless connector.^(1,2,3,4)

We have designed **vadsite** to meet these standards and furthermore, our clinical performance studies* provide the evidence to support our claim that **vadsite** meets these standards.

What do the global opinion leaders recommend?

- A needleless connector that is supported by microbial ingress testing data.⁽¹⁾
- A split septum needleless connector is associated with a lower incidence of CRBSI compared to a mechanical valve needleless connector.^(2,4)
- A needleless connector with a smooth external septum surface with few, if any, gaps, that can be more thoroughly disinfected.⁽³⁾
- A tight seal between the septum and the needleless connector housing to reduce or eliminate space for contamination to occur and biofilm to develop.⁽³⁾
- A needleless connector with a direct, that is, straight fluid pathway that facilitates adequate flushing and reduces the internal surface for biofilm development.^(3,4)
- A needleless connector with the most direct and least tortuous fluid pathway, with preferably no moving parts to reduce the potential risk of CRBSI.⁽³⁾
- A needleless connector with little or no dead space in the fluid pathway minimises the surfaces that infusates can contaminate and where biofilm can develop.⁽³⁾
- A needleless connector that does not require a clamping sequence. Or, alternatively, use only one needleless connector type that requires a specific clamp-disconnection sequence (e.g., all negative pressure, all positive pressure or all neutral pressure) throughout the healthcare facility or system and insure that all Healthcare Workers understand and are well trained in this clamp-disconnection sequence.⁽³⁾
- A transparent needleless connector is preferable to one that is opaque.⁽³⁾

* Refer to the back page for more details.

vadsite's features and benefits

• Transparent ^(3,8)

vadsite has a transparent housing and transparent silicone membrane that allows the health worker to see the fluid pathway and confirm complete flushing.

• Low Priming Volume ⁽³⁾

Suitable for clinical applications where very low priming volumes and dead space are required. Vadsite has a priming volume of just 0.07ml.

• Direct Fluid Pathway ⁽³⁾

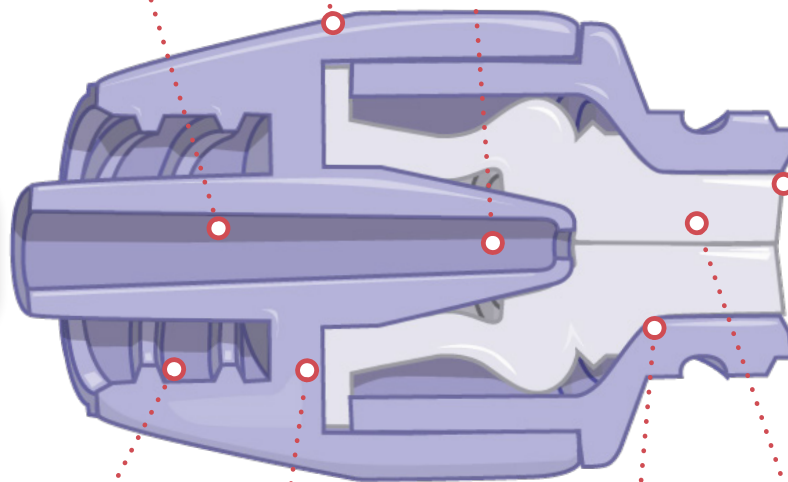
Proven 'flushable' due to its straight fluid pathway. We can prove that macro and microscopic particles, for example blood, can be successfully flushed from the device.

• Cleanable Membrane ^(1,3,6)

Proven 'cleanable' using the latest disinfectants. vadsite has a smooth easy to swab membrane that fits very tightly into the device's housing.

• MRI Compatible MR

vadsite does not contain any metal part and is therefore MRI safe.



• Effective Microbial Barrier ^(1,2,3,4,5,6)

vadsite is supported by numerous microbiological studies which conclude that microbial ingress does not occur.

• Universal Male Luer Compatibility

Any ISO standard male Luer can be connected to vadsite including those with a very small internal diameter.

• Split Septum Technology ^(2,3)

Split septum needleless connectors have demonstrated a lower incidence of CRBSI than other designs of connector.

• High Pressure Compatibility (CT-Rated)⁽⁷⁾

vadsite is approved for use with power injectors.

- Maximum pressure resistance: 350 psi
- Maximum flow rate: > 10 mL/s

References

1. Food and Drug Administration Agency (FDA), 'Guidance for Industry and FDA staff': Pre-market notification submissions, Microbial Ingress Testing, section 8, page 9, July 11th 2008.
2. Centre for Disease Control, 'Guidelines for the Prevention of Intravascular Catheter-Related Infections', Needleless Intravascular Catheter Systems, page 19, No.6. 2011.
3. William R. Jarvis, MD, 'Choosing the Best Design for Intravenous Needleless Connectors to Prevent Bloodstream Infections'. Infection Control Today, July 28th, 2010.
4. The Infusion Nurses Society, Infusion Nurses Standards of Practice; page S32, section 27, Practice Criteria A & B, 2011.
5. Evaluation of Vadsite Microbial Integrity, CARSO Laboratoire Santé, Lyon, April 2009.
6. Evaluation of Disinfection Procedures, CARSO Laboratoire Santé, Lyon, February 2011.
7. CT Pressure Testing, Laboratoire Central d'Essais, Report RE12210, Vygon SA France, August 2012.
8. Blood Clearing Analysis, 5ml Flush GLP Report, report 201301574 Rev 01, Nelson Laboratories, USA, 15th April 2013.

A summary of our clinical performance studies are detailed in the '[vadsite® Clinical Performance Studies brochure](#)'. The full protocols and results are available in '[The vadsite® Electronic Handbook](#)'. Please contact us directly or request copies directly from your local Sales Executive.

Technical specifications	
Maximum number of activations	360
Flow rate at gravity	170mL/min
High pressure flow rate	> 10 mL/s
Priming volume	0.07 mL
MRI compatible	Yes
Blood compatible	Yes
Lipids resistant	Yes
Latex-free	Yes
Alcohol resistant	Yes
Chlorhexidine resistant	Yes

- Do not put a cap, plug or obturator of any sort on **vadsite** other than a disinfecting cap.
- Do not use a needle with **vadsite**.
- Remember to clean **vadsite** prior to access.

CRITICAL CARE

For further information, please contact: questions@vygon.com

The specifications given in this brochure are for information only and are not, under any circumstances, of a contractual nature.

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