

Supplier Info Card – Viant Costa Rica, S.A.

Facility Address: Viant Costa Rica, S.A. Parque Zona Franca, Metropolitana Edificio 2C Website: www.viantmedical.com											
Facility Information Building 1: Total square feet: 31,453		Facility Information Building 2: Total square feet: 29,419									
Facility Information Building 3: Total square feet: 35,837		Facility Information Building 4: Total square feet: 81,439									
ISO Class 8 Controlled Environments: B1: 15,000 square feet B2: 7,500 square feet B4: 17,000 square feet ISO Class 7 Controlled Environment: B3: 7,500 square feet											
Company Profile: This company was previously known as “MedPlast Medical Costa Rica S.A.” The Costa Rica facility is a medical device contract manufacturing location with a Quality System certified to ISO 13485:2016, with facilities registered with the FDA.											
Total Employees: 670 Manufacturing: 550		Quality Department: 80 Sterilization is not performed on site.									
Primary Customer Contact: Site Director: Kevin Quiros Email: kevin.quiros@viantmedical.com		Phone: 506-2239-9298, ext. 6401 Fax: n/a									
A/R Contact: Luis Arce: Finance Manager Email: luis.arce@viantmedical.com		Phone: 506-2239-9298, ext. 6456 Fax: n/a									
Additional contacts: <table border="0" style="width:100%"> <tr> <td style="width:33%">Director of Quality</td> <td style="width:33%">Juan Jose Fuentes</td> <td style="width:33%">juan.fuentes@viantmedical.com</td> </tr> <tr> <td>Director of Manufacturing</td> <td>Yoryanela Perez</td> <td>yoryanela.perez@viantmedical.com</td> </tr> <tr> <td>Director of Engineering</td> <td>German Wauters</td> <td>german.wauters@viantmedical.com</td> </tr> </table>			Director of Quality	Juan Jose Fuentes	juan.fuentes@viantmedical.com	Director of Manufacturing	Yoryanela Perez	yoryanela.perez@viantmedical.com	Director of Engineering	German Wauters	german.wauters@viantmedical.com
Director of Quality	Juan Jose Fuentes	juan.fuentes@viantmedical.com									
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Juridica number: 3-101-365187											
Quality Management System: FDA Registration Number: 3005173255 ISO Registration: 13485 Certified – Quality System Registrations will remain intact and will be updated with name changes in accordance with our registrar’s standard procedures.											
Customer Quality System Audits - This information card is intended to be used by our customers as a “letter to file” that will allow previously conducted Quality System audits to remain valid until the next scheduled audit.											
Additional Facility Information: <ul style="list-style-type: none"> • 100% of our business is in the medical device market. • We do capacity planning for product development services. • We have an ERP system in place. • Customer-supplied documents are controlled. • Customer-supplied fixtures and gauges are controlled. • Customer-supplied materials are controlled. • We require our suppliers to sign a confidentiality agreement. • Our Business Excellence group leads Lean initiatives. • We monitor our facility. • Subcontractors provide services such as sterilization, component fabrication, testing labs, etc. 											