

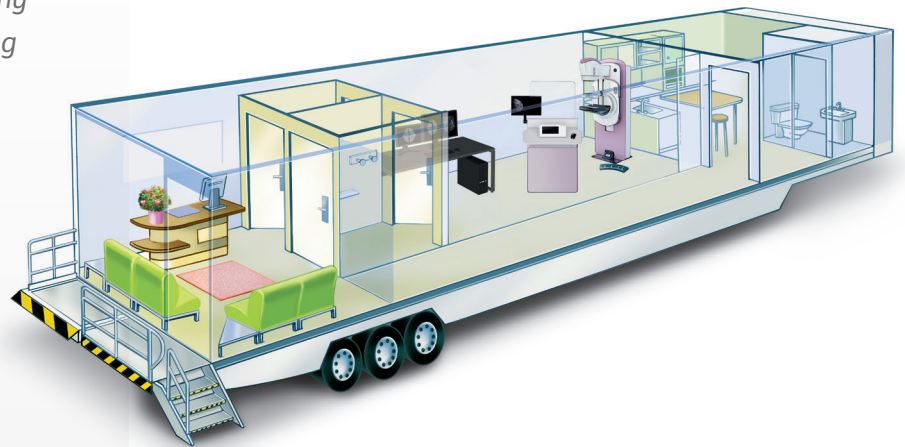
Take patient experience to the women



Convenient, Accessible Mammograms with Senographe Pristina™ Mobile

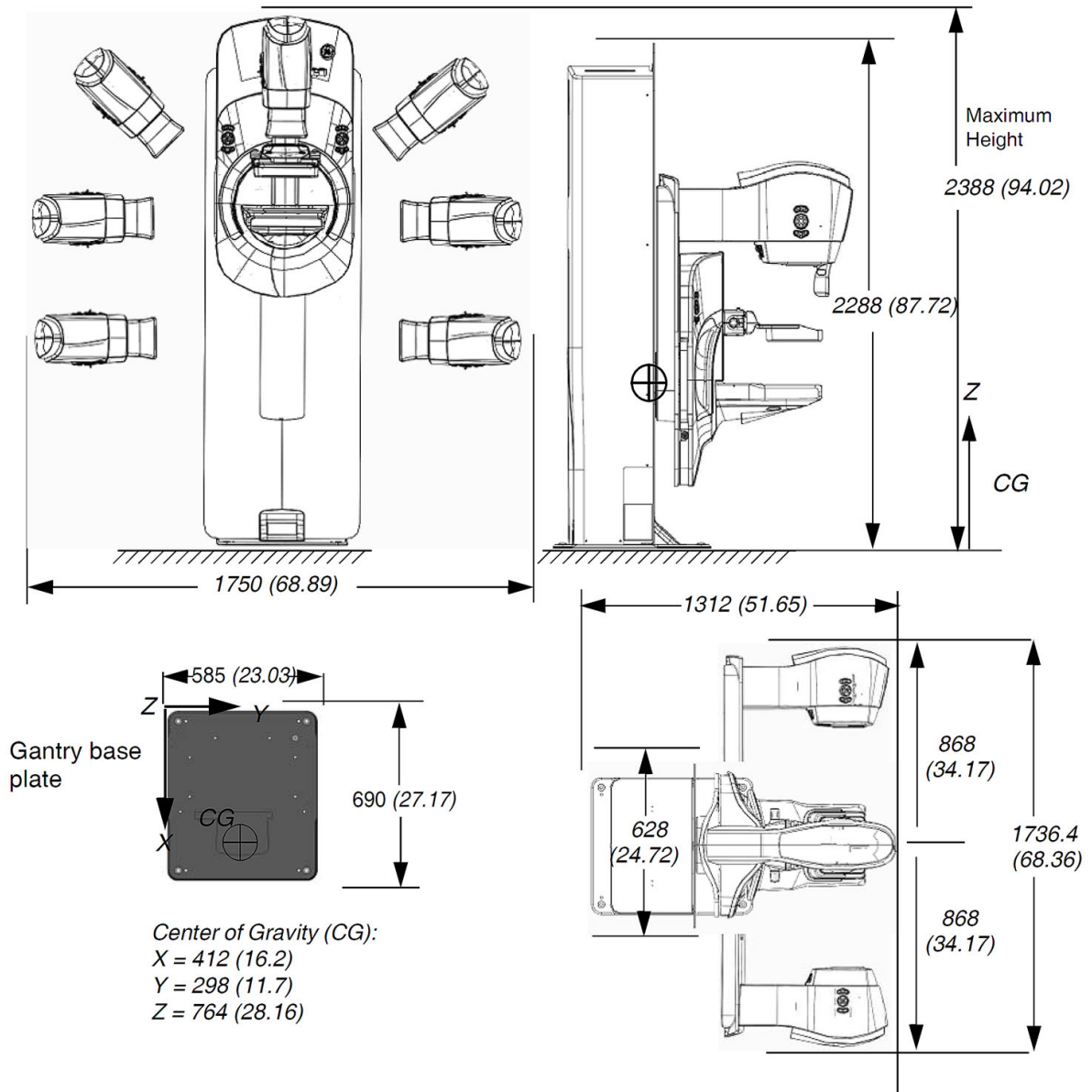
Women avoid mammograms for many reasons, anxiety and discomfort being high on that list. But another reason is **ease of access**. Meet your patients locally through patient outreach programs, work, schools, shopping centers by taking mammography to them – and reshape their mammography experience.

Bring the **patient experience and clinical benefits** to your broader community by installing Senographe Pristina on a mobile unit.



- Benefit from **Reliable and Stable** mammography platform. Secure the acquired cases until transferred
- **Experience clinical benefits** from Senographe Pristina with exceptional DBT image quality at the same low dose as a standard 2D mammography¹. Review cases in standard reading conditions with Seno Iris review
- Leverage **convenience of a better** patient experience², including Pristina Dueta™, the patient-assisted compression device (optional purchase) and **market** your new mobile mammogram option

All measurements shown as: mm (inches)



Detailed installation requirements and specifications available with purchase contract.

(1). Superior diagnostic accuracy demonstrated in a reader study comparing the ROC AUC of GE screening protocol (V-Preview + 3D CC/MLO with 3D in STD mode) to that of 2D FFDM alone. V-Preview is the 2D synthesized image generated by GE Seno Iris mammography software from GE DBT images. FDA PMA P130020 <http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpma/pma.cfm?id=P130020>.

(2). 83% of the patient rated their experience better than with previous system. IPSOS Patient Satisfaction Study sponsored by GE Healthcare, conducted with 315 patients across 2 sites in Europe, February 2017

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