Global Access Initiative is our solution to mitigating the burden of viral diseases in areas of high prevalence. This cost-effective and financially sustainable program aims to transform the world of diagnostics by improving access to critical infectious disease testing.

All-inclusive testing:
No upfront cost or capital expenditure.

$12.00 per patient test includes:

- Instrument placement†
- Reagents and consumables
- Service and maintenance
- Freight and logistics
- Replacement tests‡

Aptima® assays on the Panther® system.

As part of our commitment to the 2030 UNAIDS 95-95-95 goals; the Global Access Initiative includes HIV-1 testing and other viral targets. Qualified assays include:

- **Aptima® HIV-1 Quant Dx Assay**
  aids in the diagnosis, confirmation and clinical management of HIV-1 infection. WHO prequalified.

- **Aptima® HCV Quant Dx Assay**
  aids in the diagnosis, confirmation and clinical management of HCV infection.

- **Aptima® HBV Quant Assay**
  aids in the clinical management of HBV infection.

- **Aptima® HPV Assay**
  detects 14 high-risk HPV types.

- **Aptima® SARS-CoV-2 Assay**
  detects SARS-CoV-2 infection.

Existing sources do not include ancillary costs & sample tubes associated with sample collection, sample transport, labor costs, time, laboratory infrastructure, generic laboratory supplies & equipment, collection tubes, disposable gloves, bleach, bleach bottles, bleach enhancer and waste bottles, etc. Freight & logistics, general administration and overhead costs.

† Applies to existing instrument footprint subject to review on a case-by-case basis.
‡ Free-of-charge replacement of tests that fail due to documented instrument errors (includes one set of controls and calibrators per kit).
Expand testing in your space.

Infrastructure Requirements

To participate in the Hologic Global Access Initiative, laboratories must meet certain infrastructure requirements.

Physical Space
Sufficient space to accommodate the Panther system: 122 cm (w) x 81.5 cm (d) x 175 cm (h). Sufficient space to accommodate a rocker, centrifuge and cold storage for additional samples and reagents.

Electrical Requirements
An electrical connection that supports up to a 1400 W draw, with a dedicated 20-amp circuit.‡ A backup generator capable of supporting all equipment.

Air Conditioning
Central or wall-mounted air conditioning with adequate capacity to maintain a temperature between 15°C to 30°C and relative humidity from 20%-85%.

Sink
A sink or washbasin for liquid-waste disposal.

Waste Management
Liquid waste is nonhazardous and does not require special handling.§

Refrigerator
One or more refrigerator(s) with temperature control, maintained at 2°C to 8°C, to store samples and reagents.

Freezer
One or more freezer(s) with temperature control, maintained at -15°C to -35°C, to store calibrators and controls.

Testing Volume Requirements

To be eligible for the Hologic Global Access Initiative, the purchasing country or organization must share a site-specific forecast for any new Panther system placement, which anticipates:

Processing volumes that contribute to a national average of 30,000 patient samples per eligible instrument per year within 12 months of placement.§

Eligible Countries**

Angola
Benin
Botswana
Burkina Faso
Burundi
Cambodia
Cameroon
Cape Verde
Central African Republic
Comoros
Congo
Democratic Rep. of the Congo (DRC)
Egypt
Eswatini
Ethiopia
Gambia
Ghana
Guinea
Guinea-Bissau
Haiti
India
Indonesia
Ivory Coast
Jamaica
Kenya
Laos
Lesotho
Liberia
Malawi
Malaysia
Mali
Mauritania
Mongolia
Mozambique
Myanmar
Namibia
Nepal
Nigeria
Pakistan
Papua New Guinea
Rwanda
South Africa
Sri Lanka
Tanzania
Timor-Leste
Togo
Uganda
Vietnam
Zambia
Zimbabwe

‡ Hologic will provide an uninterruptible power supply and surge protector and make necessary electrical modifications during installation.
§ Inclusive of all eligible assays and sample types.
¶ List of eligible countries is for initiative participation. Not all countries are available in each of the countries listed.