

# Special Offer: Mix & Match

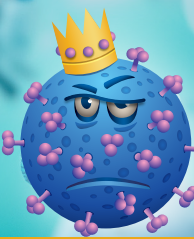
## Rapid Tests for Common Respiratory Infections



SNIFF



FLORA



SPIKE

### Take-A-Breath!

We have you covered with a broad range of rapid tests for respiratory infections.

- OSOM® COVID-19 Antigen Rapid Test (cat. no. 1066-40)
- OSOM® Ultra Plus Flu A&B Test (cat. no. 1032)
- OSOM® Flu SARS-CoV-2 Combo Test (cat. no. 1080)
  - Five kits (of any combination) must be purchased on one invoice to qualify
  - Offer valid on kits purchased July 1, 2024 - September 30, 2024 only

Get 6 Kits  
for the  
price of 5!

To receive your kits, submit redemptions online at [osompromos.com](https://osompromos.com)

Proof of purchase required

All redemptions must be received by October 31, 2024

- **Reliable:** Proven clinical performance for better patient outcomes.
- **Rapid:** Results in 15 minutes or less at the point-of-care for improved patient management.
- **Easy-to-use:** Can be performed in a CLIA-waived setting, without the need for specialized equipment.

Questions? Email us at [osomrapidtests@sekisui-dx.com](mailto:osomrapidtests@sekisui-dx.com).

Please allow up to 90 days to receive your kits. For more product information go to [osomtests.com](https://osomtests.com).



The OSOM COVID-19 Antigen Rapid Test and OSOM Flu SARS-CoV-2 Combo Test have not been FDA cleared or approved. They are authorized by FDA under an EUA for use by authorized laboratories. The OSOM COVID-19 Antigen Rapid Test has been authorized only for the detection of SARS-CoV-2 antigen, not for any other viruses or pathogens. The OSOM Flu SARS-CoV-2 Combo Test has been authorized only for the detection of proteins from SARS-CoV-2, influenza A and influenza B, not for any other viruses or pathogens. The use of these products only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. §360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.

SEKISUI Diagnostics reserves the right to change, modify, or discontinue this promotion at its discretion. Please note that the value of the special offer(s) the Customer may receive from the manufacturer under this program is a discount or other reduction in price to Customer under Section 1128B(b)(3)(A) of the Social Security Act (42 U.S.C. 1320a-7b(b)(3)(a)) and 42 C.F.R. 1001.952(h). Accordingly, Customer shall disclose this and any other discounts or other reductions in price received under this program under any state or federal program which provides cost or charge-based reimbursement to the Customer for the products and services purchased under this program. Valid for U.S. end users only, excluding Puerto Rico. The redeemed kit must be of equal or lesser value of the average price from the five purchased kits.

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**SEKISUI**  
DIAGNOSTICS

Because every result matters™