The Environmental Protection Agency (EPA) hereby grants public health exemptions under the provisions of section 18 of the Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA) as amended, to the United States Center for Disease Control and Prevention (CDC), for uses of seven antimicrobial products (comprised of six active ingredients), as listed in this document, on hard, nonporous surfaces in healthcare settings in the United States for disinfection from *Candida auris*. These exemptions are subject to the conditions and restrictions specified in the application submitted to EPA as well as the following conditions and restrictions.

1) The CDC is responsible for ensuring that all provisions of these public health exemptions are met. CDC is also responsible for providing information as specified in 40 CFR §166.32(b). Accordingly, a final report summarizing the results of this program must be submitted to the EPA Headquarters and the appropriate EPA Regional Office(s) offices within 6 months following the expiration of this public health exemption. Additionally, in accordance with 40 CFR §166.32(a), these offices shall be immediately informed of any adverse effects resulting from the use of these chemicals in connection with these exemptions. Refer to file symbols 19-DO-03, -04, -05, -06, -07, and -08 in any future correspondence regarding these exemptions.

2) The uses may take place at healthcare facilities in the United States, to disinfect hard, nonporous, non-food/feed-contact surfaces, including facilities, furnishings, and equipment, that may be contaminated with *Candida auris*, when registered alternatives are not suitable for use, particularly where sensitive or compromised individuals are present.
3) The following products may be used under these exemptions:

a) Oxivir TB Spray (EPA Reg# 70627-56); containing 0.5% hydrogen peroxide, manufactured by Diversy
b) Oxivir TB Wipes (EPA Reg# 70627-60); containing 0.5% hydrogen peroxide, manufactured by Diversy
c) Hydrogen Peroxide Disinfectant Spray (EPA Reg# 67619-24); containing 1.4% hydrogen peroxide, manufactured by Clorox
d) Hydrogen Peroxide Disinfectant Wipes (EPA Reg# 67619-25); containing 1.4% hydrogen peroxide, manufactured by Clorox
e) PDI Sani Prime Spray (EPA Reg# 9480-10); containing 0.61% Didecyl dimethyl Ammonium chloride, 28.7% isopropanol, and 27.3% ethanol; manufactured by PDI
f) PDI Sani-Cloth Prime (EPA Reg# 9480-12); containing 0.61% didecyl dimethyl ammonium chloride, 28.7% isopropanol, and 27.3% ethanol; manufactured by PDI
g) PDI Super Sani-cloth (EPA Reg# 9480-4); containing 0.5% quaternary ammonia compounds (0.25% n-alkyl dimethyl ethylbenzyl ammonium chlorides, 0.25% n-alkyl dimethyl benzyl ammonium chlorides), and 55% isopropanol; manufactured by PDI

4) Follow all use directions, restrictions and precautions on the registered product labels for germicidal activity against Candida albicans when applicable. When referencing germicidal activity against C. albicans is not applicable, apply product according to existing label usage for fungicidal activity, including any requirements for precleaning of surfaces or personal protective equipment.

5) No treatments are permitted under this authorization on food or feed items, or where food or feed is present.

6) This public health exemption expires one year from date of authorization, and a final report is due within six months after expiration, as given above.

7) If any of the requested products become registered for C. auris during the duration of these exemptions, they should be used under their registrations and their exemption will no longer be valid.

If you have any questions regarding the authorization of this quarantine exemption program, please contact Emergency Response Team member Andrea Conrath at (703) 308-9356; conrath.andrea@epa.gov.

Richard P. Keigwin, Jr., Director
Office of Pesticide Programs

Date: 10/11/2019