UNIVERSITY OF MARYLAND BALTIMORE
PROCESS FOR RESUMPTION OF HUMAN RESEARCH

FOLLOW THESE STEPS:

1. Principal Investigators will have prioritized their research protocols (in Phase 0) in the order that they wish them to be considered for resumption.

2. Complete this UMB Environmental Health and Safety Checklist (below) for Resuming Clinical Research FOR EACH currently approved (UMB or external IRB) human research protocol the Principal Investigator want to resume. This tool can be found at https://www.hrpo.umaryland.edu and the UMB COVID-19 website https://www.umaryland.edu/coronavirus/

3. Principal Investigators should ensure that their individual plans are consistent with their School’s and Department’s Operations Assessment Tool. This tool can be found at https://www.hrpo.umaryland.edu and the UMB COVID-19 website https://www.umaryland.edu/coronavirus/

4. All requests for resumption of research must be submitted under the appropriate Stage. Any requests that are submitted during an incorrect Stage will be denied and must be re-submitted during the appropriate Stage for consideration.

5. Submit BOTH this Checklist and the Operations Assessment Tool through the Reportable New Information (RNI) pathway in CICERO.

6. It is mandatory for ONLY the Principal Investigator to submit the RNI in CICERO for each study. RNI's from team members will not be accepted.

7. The Checklist and the Operations Assessment Tool will be reviewed by institutional representatives for appropriateness.

8. Acknowledgement of your submission through CICERO RNI pathway will be the documentation to proceed with resumption of research.

9. Do not proceed with research resumption until such time as you receive the acknowledgment in CICERO.

For questions regarding completion of these forms please contact Dr. Julie Doherty at jdoherty@umaryland.edu