### Things to think about…

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<th>Yes</th>
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| 1. | Before you begin to plan your study contact a WashU IRB Partner for consultation and advice prior to IRB submission. HRPO SWAT: 314-747-6800.  
  - HRPO website [https://hrpo.wustl.edu/](https://hrpo.wustl.edu/)  
  - myIRB research application program [https://myirb.wusm.wustl.edu/](https://myirb.wusm.wustl.edu/) | ☐ | ☐ | ☐ |
| 2. | Identify the international site’s local regulatory requirements and reach out to your HRPO Partner to see whether or not they are already aware of your international site’s regulatory requirements. | ☐ | ☐ | ☐ |
| 3. | If local IRB or Ethical Committee review [IS](#) required:  
  - Identify the processes needed to obtain the review.  
  - Have you obtained the appropriate documentation (Letter of Approval)?  
  - Please note: depending on the location, this review may take the form of a letter of approval from an IRB or research ethics committee, national regulatory office, local university department sponsoring the research, institutional oversight committee. | ☐ | ☐ | ☐ |
| 4. | If local IRB or ethical committee review is [NOT](#) required or is not available:  
  - Identify a Cultural Reviewer/Expert who can provide you with documentation of cultural appropriateness. This expert cannot be part of the study.  
  - Identify the appropriate Regulatory Official or individual who can provide you with documentation that local IRB or ethics review is not required OR documentation of what the local requirements for conduct of human subjects research are. | ☐ | ☐ | ☐ |
| 5. | Do your research documents need to be translated? Be sure to document the names and qualifications of translators for data collection instruments, informed consents etc. | ☐ | ☐ | ☐ |
| 6. | Are specific international site permissions required?  
  - Do you know the processes for obtaining these permissions? Or when permission(s) cannot be obtained in advance, are you familiar with the procedures to acquire permission(s) prior to initiating research activities?  
    - Note: some international sites may not allow you to obtain these permissions until you are in the country when the research commences. In these cases, it is good to know the process so you can communicate that to the WashU IRB.  
  - Once received, do you have documentation of the permission(s)? | ☐ | ☐ | ☐ |
| 7. | Are there country-specific licenses, permits or other authorizations necessary for the procedures to be performed at the international site? For example, licensure for clinical procedures or permits for drugs, devices, or technology being brought into the foreign country. | ☐ | ☐ | ☐ |
| 8. | Do you want to compensate or give class credit to your research participants?  
  - If yes, are there any local laws or regulations that prevent you from doing this?  
  - What is a culturally appropriate way to compensate them or offer extra credit? | ☐ | ☐ | ☐ |
| 9. | Are you going to ask local individuals (non-Washington University faculty, staff or students) to assist with study procedures, recruitment, and informed consent in the international setting?  
  - If yes, be sure to collect & document their names, qualifications, and protocol training. This is only required when the international sites do not have a local IRB or equivalent ethics committee approval (i.e. there is no local IRB or ethics committee and you wish for WashU IRB to serve as the IRB for the local research staff).  
  - Be sure to clearly describe their role and identify them as non-WashU research staff in your IRB application | ☐ | ☐ | ☐ |
| 10. | Are there country-specific laws or regulations about data collection and/or data transportation?  
  - Are there any country-specific laws or regulations about storing consent forms, research data/documentation and/or any identifiable data?  
  - Have you reviewed WashU’s guidance on electronic storage of research documents? [https://research.wustl.edu/electronic-storage-research-study-documents/](https://research.wustl.edu/electronic-storage-research-study-documents/)  
### International Research Prep Work Checklist – Prior to Creating IRB Application

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<td><strong>11.</strong> If the research is federally funded, have you provided the FWA number assigned to the foreign site?</td>
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| **12.** Is your consent process culturally appropriate?  
  - Consider discussing the consent process with your cultural expert.  
  - Contact your WashU HRPO/IRB Partner for advice and guidance on different consent processes that may be used.  
  - Build in flexibility | ☐  | ☐  | ☐  |
| **13.** Have you built in flexibility into your research protocol?  
  - For example, if you plan to interview participants in several places and not just a single location, be sure to describe all options.  
  - Use the term “may” if one or more study procedure could be used. Be sure to ask your IRB Partner for suggestions if the “normal” process makes implementing the research too difficult or culturally inappropriate. | ☐  | ☐  | ☐  |
| **14.** If you are student, have you identified your WashU Faculty Sponsor?  
  - Please review the following with your Faculty Sponsor:  
    - All research team members, including your Faculty Sponsor, must have CITI human subjects training in order to be added to the research team in myIRB.  
    - This can cause delays if you or your Faculty Sponsor has not completed this training.  
    - IF non-WashU individuals will be participating in the research, be sure to work with you HRPO partner to develop appropriate training for the participating non-WashU staff.  
    - The Faculty Sponsor is ultimately responsible for overseeing the research you are conducting. It is highly important that you make arrangements (and document them) to review your research records with your Faculty Sponsor. | ☐  | ☐  | ☐  |
| **15.** Have you thought about your research documentation?  
  - It is very important that you document what you do, when you do it, and with whom it is done with. It is also equally as important to document when something cannot be done and why.  
    - Always remember…If it’s not documented, then it did not happen.  
    - The Human Subjects Research Website provides forms and templates that you can use to document your research: [https://research.wustl.edu/topics/human-subjects-research/](https://research.wustl.edu/topics/human-subjects-research/)  
    - Contact the Human Research Quality Assurance Program for additional resources or research documentation support. [HRQA@wustl.edu](mailto:HRQA@wustl.edu) or +1 314-747-5525 | ☐  | ☐  | ☐  |
| **16.** Have you consulted the WashU policies about taking data or research materials with you if/when you may permanently leave WashU for a job at another university? | ☐  | ☐  | ☐  |