## University of Kentucky Office of Research Integrity Exemption Categories Tool

- Subpart B: Studies Involving Pregnant Women, Fetuses & Neonates are Eligible for Exempt Under All 8 Categories
- Subpart C: Exemptions Do Not Apply to Research Involving Prisoners <u>Except</u> "for Research Aimed at Involving a Broader Subject Population that <u>Only Incidentally</u> Includes Prisoners"
- Subpart D: Children are allowed in categories 1,4,5,6,7, & 8; Limitations & Exclusion of Children in Category 2 & 3

Category	New Citation	Exemption Category Description	Limited IRB Review	Conditions/Allowances/Limitations
1	104(d)(1)	Research in Established or Commonly Accepted Education Settings that Involves Normal Educational Practices	N/A	Not Likely to Adversely Impact Students' Opportunity to Learn or Assessment of Educators Providing Instruction
2	104(d)(2)	Research only includes interactions involving Educational Tests, Surveys, Interviews, Public Observation if at least ONE of the following criteria met:	N/A	Data Collection Only; May include visual or auditory recording; May NOT include Intervention Only includes interactions
		<ul> <li>(i) Recorded information cannot readily identify the subject (directly or indirectly/linked)</li> </ul>	N/A	Surveys & Interviews: No Children; Educational Tests or Observation of Public Behavior: Can Only include Children When Investigators Do Not Participate in Activities being Administered/Observed
		<ul> <li>(ii) Any disclosure of responses outside of the research would NOT reasonably place subject at risk (criminal, civil liability, financial, employability, educational advancement, reputation)</li> </ul>	N/A	Surveys & Interviews: No Children; Educational Tests or Observation of Public Behavior: Can Only include Children When Investigators Do Not Participate in Activities being Administered/Observed
		<ul> <li>(iii) Information is recorded with identifiers or code linked to identifiers &amp; IRB conducts Limited Review</li> </ul>	Privacy and Confidentiality Review	NO Children
3	104(d)(3)(i)	Research involving Benign Behavioral Interventions (BBI) through verbal, written responses, (including data entry or audiovisual recording) from adult subject who prospectively agrees and ONE of following met:	N/A	NO Children; May Not include Medical Interventions; Subject prospectively agrees; BBI must be:
		A. Recorded information cannot readily identify the subject (directly or indirectly/linked)	N/A	<ul> <li>Brief in Duration</li> <li>Painless/Harmless</li> <li>Not Physically Invasive</li> <li>Not Likely to Have a Significant Adverse</li> </ul>
		<ul> <li>B. Any disclosure of responses outside of the research would NOT reasonably place subject at risk (criminal, civil liability, financial, employability, educational advancement, reputation)</li> </ul>	N/A	Lasting Impact on Subjects <ul> <li>Unlikely that Subjects Will Find</li> <li>Interventions Offensive or</li> <li>Embarrassing</li> </ul> No deception unless participant is informed in
		C. Information is recorded with identifiers & IRB conducts Limited Review	Privacy and Confidentiality Review	the prospective agreement that he/she will be unaware of or misled regarding the true nature or purpose of the research

Category	New Citation	Exemption Category Description	Limited IRB Review	Conditions/Allowances/Limitations
4	104(d)(4)	Secondary Research for Which Consent is Not Required: Use of Identifiable Information or Identifiable Biospecimen that have been or will be collected for some other 'primary' or 'initial' activity, if ONE of following criteria met:		No Primary Collection from subjects for the research; Allows Both <u>Retrospective and Prospective</u> <u>Secondary Use</u>
		(i) Biospecimens or Information is Publicly Available	N/A	Must be publicly available (e.g., commercially available specimen or open access data). May also qualify as not human research (NHR).
		<ul> <li>(ii) Information recorded so subject cannot readily be identified (directly or indirectly/linked); Investigator does not contact subjects and will not re-identify the subjects</li> </ul>	N/A	PI does not contact; Will not re-identify
		<ul> <li>(iii) Collection and Analysis involving Investigators Use of Identifiable Health Information when use is regulated by HIPAA "health care operations" or "research" or "public health activities and purposes"</li> </ul>	N/A	HIPAA regulations still apply; HIPAA protections include authorization or waiver of authorization; Does not include Biospecimens (only PHI); Only covers "investigator's use"; does not indicate that sharing is permitted under this exemption.
		(iv) Research information collected by or on behalf of federal government using government generated or collected information obtained for non-research activities	N/A	If research generates identifiable private information it is subject to specified federal privacy laws (see iv for list)
5	104(d)(5)	Research and demonstration projects supported by a Federal Agency/Dept. AND Designed to studyimprove public benefit or service programs.	N/A	Must be posted on a Federal Web Site
6	104(d)(6)	Taste and Food Quality – no change	N/A	Wholesome food without additives;
	104(-1)(-7)		- If there is a	ingredient level and use found to be safe (see ii)
7*	104(d)(7)	Storage or Maintenance of Identifiable Private Information or Identifiable Biospecimens for Secondary Research For Which Broad Consent Is Required	- If there is a change made for research purposes in the way material stored or maintained, Privacy and confidentiality review	All requirements for Broad Consent Met; -Broad consent is obtained Documented or documentation waived MUST TRACK REFUSALS —as the IRB may not waive consent for use of identifiable material for any individual who refuses
8*	104(d)(8)	Secondary research involving use of Identifiable Private Information or Identifiable Biospecimens for Which Broad Consent was Required *UK NOT USING	-Privacy and confidentiality review & -research is within the scope of the broad consent & -PI does not plan to return research results	Privacy and Confidentiality protections adequate; Broad consent was obtained; Documented or documentation waived No plan to return research results; MUST TRACK REFUSALS as the IRB may not waive consent for use of identifiable material for any individual who refuses