

BIBLIOGRAFÍA

Ética de la investigación biomédica

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Esta selección bibliográfica ha sido realizada en julio de 2007 mediante una búsqueda en la base de datos electrónica Philosopher's Index, cruzando las siguientes palabras clave (opción "cualquier campo"):

[("ETHICS-" in DE,PS) and (((("EXPERIMENTATION-" in DE,PS) or ((("MEDICAL-RESEARCH") in DE,PS or (research ethics) or ((("CLINICAL-TRIAL") in DE,PS))) and (research ethics)]

or

[("BIOETHICS-" in DE,PS) and (((("EXPERIMENTATION-" in DE,PS) or ((("MEDICAL-RESEARCH") in DE,PS) or (research ethics) or ((("CLINICAL-TRIAL") in DE,PS))].

Un total de 183 entradas fueron halladas. De ellas, sólo se seleccionaron las publicaciones posteriores a 1996 relacionadas con 5 subtemas: consentimiento informado, comités de ensayos clínicos, conflicto de intereses, selección de sujetos de investigación y poblaciones vulnerables e investigación biomédica en países en vías de desarrollo. Esta selección fue completada con referencias que no aparecieron en la primera búsqueda, por tratarse generalmente de fuentes no filosóficas. Para ello se realizó una búsqueda complementaria (similares palabras clave) en las bases de datos del CSIC (sección Filosofía) y PubMed, de Medicina. Debido a la excesiva cantidad de referencias encontradas, sólo

se seleccionaron aquellas que, de acuerdo con el criterio del compilador, parecieran más relevantes. En la bibliografía aparecen por orden alfabético las entradas correspondientes con cada uno de los campos temáticos citados.

Al final de la selección, se incluyen dos apartados: "Monografías", que contiene referencias encontradas a través de una búsqueda en Philosopher's Index (límite "monografías"), y en los sitios de Internet de bioethics.net y Amazon; y "Referencias clásicas", -apartado que incluye algunos textos fundacionales de la ética de la investigación biomédica.

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