

Supplemental Material

Title

Weaponized uranium and adverse health outcomes in Iraq: A systematic review

Authors

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Table S1. PECOS Statement and inclusion/exclusion criteria

Study Aspect	Inclusion Criteria	Exclusion Criteria
Population	Humans (of any age) residing in Iraq at any time between 1990-2020	Non-humans
	Children born to those individuals	Military veterans (U.S. or other nationalities)
Exposure	Chemical exposure to metallic uranium that has been introduced to the environment in Iraq via the use of conventional weapons (i.e. non-nuclear missiles, bullets, and armor)	Physical exposure to ultraviolet radiation (e.g. solar radiation)
	Chemical exposure to decay and corrosion products of weapon uranium in Iraq	Physical exposure to ionizing radiation emitted from medical radioisotopes
	Physical exposure to ionizing radiation emitted from radioisotopes in weaponized uranium or its decay or erosion products	Physical or chemical exposure to nuclear materials not originating from metallic uranium weapons used by US and coalition forces
Comparator	Individuals not exposed to uranium	Studies that document health impacts among an exposed population without comparison to a non-exposed (or lesser exposed) group or population
	Individuals exposed to lower levels	
	Can include historical controls	
Outcomes	Human health-relevant outcomes, including measures of general wellbeing, mental health, or self-rated health	Studies that measure the concentration of uranium (or decay/corrosion products) in environmental or human biological samples without measuring a health outcome
		Studies that measure radiation levels in food, water, or environmental samples without measuring a health outcome
Study Design	Observational study designs including case-control, case-report, cohort, and cross-sectional	Randomized control trials or other experimental study designs
		Reviews (including systematic reviews) that do not include or report primary research

Table S2. Full search strategy

Database	MEDLINE Ovid MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily, 1946 to April 06, 2020
Search	(exp URANIUM COMPOUNDS/ or exp URANIUM/ or exp RADIOLOGIC HEALTH/ or THORIUM/ or PLUTONIUM/ or PROTACTINIUM/ or exp THORIUM COMPOUNDS/ or (uranium* or diuranium* or triuranium* or DU or "U(VI)" or "U(IV)" or U-235 or U235 or U-238 or U238 or Qmetal* or (Q adj metal*) or depletalloy* or (deplet* adj alloy*) or uranyl or radiation* or radioactiv* or radioisotop* or radionuclide* or radiologic* or dosimet* or ((gamma* or beta* or alpha* or nuclear) adj3 (ray? or radiation? or emitt* or emission* or decay*)) or plutonium* or thorium* or protactinium*).mp.) AND (exp IRAQ/ or exp IRAQ WAR, 2003-2011/ or exp GULF WAR/ or (Iraq* or (operation* adj2 ((new adj dawn*) or freedom*)) or (gulf adj2 (war* or Arab* or Persian*)) or (operation* adj2 (desert* adj (storm* or shield*))) or mosul* or Falluja* or Al-Anbar* or alanbar* or Anbar* or Babil* or Baghdad* or Al-Basra* or Albasra* or Basra* or ((Dhi or Thi*) adj Qar) or Al-Qadisiy* or Alqadisiy* or Qadisiy* or Diyala* or Dohuk* or Erbil* or Halabja* or Karbala* or Kirkuk* or Maysan* or Al-Muthan* or almuthan* or Muthan* or Najaf* or Ninev* or Saladin* or Sulaymaniy* or Wasit* or kurd*).mp.) AND (1990:2020.(sa_year).) NOT (Animals/ not (Animals/ and Humans/))
Results	355
Database	Embase
Search	('uranium'/exp OR 'uranium derivative'/exp OR 'radiation and radiation related phenomena'/exp OR 'thorium'/exp OR 'plutonium'/exp OR 'protactinium'/exp OR 'thorium derivative'/exp OR uranium*:ab,ti OR diuranium*:ab,ti OR triuranium*:ab,ti OR du:ab,ti OR 'u(vi)':ab,ti OR 'u(iv)':ab,ti OR 'u-235':ab,ti OR u235:ab,ti OR 'u-238':ab,ti OR u238:ab,ti OR qmetal*:ab,ti OR ((deplet* NEAR/2 alloy*):ab,ti) OR uranyl:ab,ti OR radiation*:ab,ti OR radioactiv*:ab,ti OR radioisotope*:ab,ti OR radionuclide*:ab,ti OR radiological*:ab,ti OR dosimet*:ab,ti OR

	<p>plutonium*:ab,ti OR thorium*:ab,ti OR protactinium*:ab,ti OR (((gamma* OR beta* OR alpha* OR nuclear) NEAR/3 (ray* OR radiation* OR emitt* OR emission* OR decay*)):ab,ti) AND ('iraq'/exp OR 'iraqi'/exp OR 'iraqi kurdistan'/exp OR 'persian gulf'/exp OR iraq*:ab,ti OR ((operation* NEAR/2 new* NEAR/2 dawn*):ab,ti) OR ((operation* NEAR/2 freedom*):ab,ti) OR ((gulf NEAR/2 war*):ab,ti) OR ((gulf NEAR/2 arab*):ab,ti) OR ((gulf NEAR/2 persian*):ab,ti) OR ((operation* NEAR/2 desert* NEAR/2 storm):ab,ti) OR ((operation* NEAR/2 desert* NEAR/2 shield*):ab,ti) OR mosul*:ab,ti OR falluja*:ab,ti OR 'al-anbar*':ab,ti OR alanbar*:ab,ti OR anbar*:ab,ti OR babil*:ab,ti OR baghdad*:ab,ti OR 'al-basra*':ab,ti OR albasra*:ab,ti OR basra*:ab,ti OR ((dhi* NEAR/2 qar):ab,ti) OR ((thi* NEAR/2 qar):ab,ti) OR 'al-qadisy*':ab,ti OR alqadisy*:ab,ti OR qadisy*:ab,ti OR diyala*:ab,ti OR dohuk*:ab,ti OR erbil*:ab,ti OR halabja*:ab,ti OR karbala*:ab,ti OR kirkuk*:ab,ti OR maysan*:ab,ti OR 'al-muthan*':ab,ti OR almuthan*:ab,ti OR najaf*:ab,ti OR ninev*:ab,ti OR saladin*:ab,ti OR sulaymaniy*:ab,ti OR wasit*:ab,ti OR kurd*:ab,ti) AND [1990-2020]/py NOT ([animals]/lim NOT [humans]/lim)</p>
Results	627
Database	PubMed
Search	<p>((iraq[mesh]) OR (iraq war, 2003-2011[mesh]) OR (gulf war[mesh]) OR Iraq*[tw] OR operation new dawn*[tw] OR operation iraqi freedom[tw] OR gulf war*[tw] OR persian gulf*[tw] OR arab gulf*[tw] OR operation desert storm*[tw] OR operation desert shield*[tw] OR mosul*[tw] OR Falluja*[tw] OR Al-Anbar*[tw] OR alanbar*[tw] OR Anbar*[tw] OR Babil*[tw] OR Baghdad*[tw] OR Al-Basra*[tw] OR Albasra*[tw] OR Basra*[tw] OR dhiqar*[tw] OR thiqar*[tw] OR dhi-qar*[tw] OR thi-qar*[tw] OR Al-Qadisiy*[tw] OR Alqadisiy*[tw] OR Qadisiy*[tw] OR Diyala*[tw] OR Dohuk*[tw] OR Erbil*[tw] OR Halabja*[tw] OR Karbala*[tw] OR Kirkuk*[tw] OR Maysan*[tw] OR Al-Muthan*[tw] OR almuthan*[tw] OR Muthan*[tw] OR Najaf*[tw] OR Ninev*[tw] OR Saladin*[tw] OR Sulaymaniy*[tw] OR Wasit*[tw] OR kurd*[tw]) AND ((uranium[mesh]) OR (uranium compounds[mesh]) OR (radiologic health[mesh]) OR (thorium[mesh]) OR (thorium compounds[mesh]) OR (plutonium[mesh]) OR (protactinium[mesh]) OR (uranium*[tw] OR uranyl*[tw] OR plutonium*[tw] OR thorium*[tw] OR protactinium*[tw] OR qmetal*[tw]</p>

	OR depletalloy*[tw] OR q-metal*[tw] OR deplete-alloy*[tw] OR radiation*[tw] OR du[tw] OR u235[tw] OR u238[tw] OR u-235[tw] OR u-238[tw] OR radiologic*[tw] OR radioactiv*[tw] OR radioisotop*[tw] OR radionuclide*[tw] OR dosimet*[tw] OR gamma-ray*[tw] OR gamma-emit*[tw] OR gamma-emission*[tw] or gamma-decay*[tw] OR beta-ray*[tw] OR beta-emit*[tw] OR beta-emission*[tw] OR beta-decay*[tw] OR alpha-ray*[tw] or alpha-emit*[tw] OR alpha-emission*[tw] OR alpha-decay*[tw] OR nuclear-ray*[tw] or nuclear-emitt*[tw] or nuclear-emission*[tw] or nuclear-decay*[tw])) NOT ("Animals"[Mesh] NOT ("Animals"[Mesh] AND "Humans"[Mesh])) AND ("1990/01/01"[PDAT] : "2020/12/31"[PDAT])
Results	317
Database	Scopus
Search	TITLE-ABS-KEY("Iraq*" OR (Gulf W/2 War*) OR ((Persia* or Arab*) W/2 Gulf) OR (operation* W/2 new* W/2 dawn*) OR (operation* W/2 freedom*) OR (operation* W/2 desert* W/2 storm*) OR (operation* W/2 desert* W/2 shield*) OR mosul* or Falluja* or *Anbar* or Babil* or Baghdad* or *Basra* or ((Dhi or Thi*) W/2 Qar) or *Qadisiy* or Diyala* or Dohuk* or Erbil* or Halabja* or Karbala* or Kirkuk* or Maysan* or *Muthan* or Najaf* or Ninev* or Saladin* or Sulaymaniy* or Wasit* or kurd*) AND TITLE-ABS-KEY(*uranium* OR thorium* OR plutonium* OR protactinium* OR DU OR "U(VI)" OR "U(IV)" OR U235 OR U-235 OR U238 OR U238 OR qmetal* OR q-metal* or depletalloy* OR (deplet* W/2 alloy*) or uranyl or radiation* or radioactiv* or radioisotop* or radionuclide* or radiologic* or dosimet* or ((gamma* or beta* or alpha* or nuclear) w/3 (ray* or radiation* or emitt* or emission* or decay*))) AND PUBYEAR > 1990
Results	1546
Database	TOXLINE

Search	((uranium OR "uranium 238" OR 7440-61-1 [rn]) OR uranyl OR plutonium* OR thorium* OR protactinium* OR qmetal* OR depletalloy* OR "q-metal" OR "deplete-alloy" OR radiation* OR du OR u235 OR u238 OR "u-235" OR "u-238" OR radiologic* OR radioactiv* OR radioisotop* OR radionuclide* OR dosimet* or "alpha-ray" or "alpha ray" or "alpha-emitter" OR "alpha emitter" OR "alpha-emission" OR "alpha emission" OR "alpha-decay" or "alpha decay" or "beta-ray" or "beta ray" or "beta-emitter" or "beta emitter" or "beta-emission" or "beta emission" or "beta-decay" or "beta-decay" or "gamma-ray" or "gamma ray" or "gamma-emitter" or "gamma emitter" or "gamma-emission" or "gamma emission" or "gamma-decay" or "gamma decay" or "nuclear-ray" or "nuclear ray" or "nuclear-emitter" or "nuclear emitter" or "nuclear-emission" or "nuclear emission" or "nuclear-decay" or "nuclear decay") AND (iraq* OR "gulf war" OR "persian gulf" OR "arab gulf" OR "desert storm" OR "desert shield" OR "operation iraqi freedom" OR "operation new dawn" OR mosul* OR falluja* OR al-anbar* OR alanbar* OR anbar* OR babel* OR baghdad* OR al-basra* OR albasra* OR basra* OR dhi-qar OR thi-qar OR dhiqar OR thiqr OR al-qadisiy* OR alqadisiy* OR qadisiy* OR diyala* OR dohuk* OR erbil* OR halabja* OR karbala* OR kirkuk* OR maysan* OR al-muthan* OR almuthan* OR muthan* OR najaf* OR ninev* OR saladin* OR sulaymaniy* OR wasit* OR kurd*) AND 1990:2020 [yr]
Results	462
Database	Iraqi Academic Scientific Journals
Search	(all:Uranium* all:or all:U-235 all:or all:U-238 all:or all:U235 all:or all:U238) Publication Year: 1990 to 2020
Results	229
Database	ProQuest Dissertations and Theses Global

Search	TI,AB,SU((uranium* OR uranyl OR plutonium* OR thorium* OR protactinium* OR qmetal* OR depletalloy* OR "q-metal" OR "deplete-alloy" OR radiation* OR du OR u235 OR u238 OR "u-235" OR "u-238" OR radiologic* OR radioactiv* OR radioisotop* OR radionuclide* OR dosimet* or "alpha-ray" or "alpha ray" or "alpha-emitter" OR "alpha emitter" OR "alpha-emission" OR "alpha emission" OR "alpha-decay" or "alpha decay" or "beta-ray" or "beta ray" or "beta-emitter" or "beta emitter" or "beta-emission" or "beta emission" or "beta-decay" or "beta-decay" or "gamma-ray" or "gamma ray" or "gamma-emitter" or "gamma emitter" or "gamma-emission" or "gamma emission" or "gamma-decay" or "gamma decay" or "nuclear-ray" or "nuclear ray" or "nuclear-emitter" or "nuclear emitter" or "nuclear-emission" or "nuclear emission" or "nuclear-decay" or "nuclear decay") AND (iraq* OR "gulf war" OR "persian gulf" OR "arab gulf" OR "desert storm" OR "desert shield" OR "operation iraqi freedom" OR "operation new dawn" OR mosul* OR falluja* OR al-anbar* OR alanbar* OR anbar* OR babel* OR baghdad* OR al-basra* OR albasra* OR basra* OR dhi-qar OR thi-qar OR dhiqar OR thiqar OR al-qadisiy* OR alqadisiy* OR qadisiy* OR diyala* OR dohuk* OR erbil* OR halabja* OR karbala* OR kirkuk* OR maysan* OR al-muthan* OR almuthan* OR muthan* OR najaf* OR ninev* OR saladin* OR sulaymaniy* OR wasit* OR kurd*)) AND YR(1990-2020)
Results	52
Database	Google Scholar
Search	Allintitle: Uranium Iraq Publication Year: 1990-2020
Results	122
Database	IAEA Scientific and Technical Publications

Search	Search: Uranium Publication Year: 1990-2020 Search: Iraq Search: Iraqi
Results	116
Database	WHO Institutional Repository for Information Sharing
Search	Iraq* AND Uranium
Results	43
Database	UNEP Knowledge Repository
Search	Iraq
Results	22

Table S3. Preferred reporting items for systematic reviews and meta-analysis (PRISMA) guidelines

Section/topic	#	Checklist item	Reported on page #
TITLE			
Title	1	Identify the report as a systematic review, meta-analysis, or both.	1
ABSTRACT			
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.	2
INTRODUCTION			
Rationale	3	Describe the rationale for the review in the context of what is already known.	3-5
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).	5
METHODS			
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.	6-7
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.	6; Supplemental Material (Table S1)
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.	6; Supplemental Material (Table S2)
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.	Supplemental Material (Table S2)
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).	6; Supplemental Material "Screening Form" and "Full text exclusion justifications"
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.	7; Supplemental Material "Data

			abstraction form”
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.	7-8; Supplemental Material Tables S6-S13
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.	7; Supplemental Material “Navigation Guide instructions for making risk of bias determinations”
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).	7
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I^2) for each meta-analysis.	No meta-analysis was conducted
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).	7-8, Supplemental Material Table S5
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	No meta-analysis was conducted
RESULTS			
Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.	9, Figure 1
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.	9-10; Supplemental Material Tables S5-S13
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).	10-15, Figures 3A & 3B, Supplemental Material Tables S14-S49
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.	9-15, Table 1, Supplemental Material Tables S5-S13
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.	No meta-analysis was conducted
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).	10-15, Table 1, Figure 3A & 3B

Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).	No meta-analysis was conducted
DISCUSSION			
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).	10-15, Table 1
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).	17
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.	16-19
FUNDING			
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.	20

Table S4. Quality of evidence grades (Schünemann et al. 2013)

Grade	Definition
High	We are very confident that the true effect lies close to that of the estimate of the effect.
Moderate	We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.
Low	Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect.
Very Low	We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect.

Table S5. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and congenital birth defects among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Alaani et al. (2012)	Case-control	Nov. 2009 – Sept. 2010	Births in Fallujah	Population based	Place of residence (Exposed = Fallujah, Unexposed = Egypt, Kuwait, and UAE, i.e. historical control)	CBD incidence rate (number of infants born with CBD per 1000 live births in population)	No statistical test	CBD incidence in Fallujah for the 11 month period was reported to be between 48-144/1000 live births. This rate is many times higher than CBD incidence reported for other (unexposed) Arab countries (Giza, Egypt 31.7, Kuwait 12.5, UAE 7.9)
Alaani et al. (2011)	Case-control	2009-2010	Parents who gave birth to children with CBD at Fallujah General Hospital	Cases = 25 Controls = 99, (Israeli) 114 (Swedish), 2 (Iraqi)	Uranium concentration in hair scalp samples (U mg·kg ⁻¹) and uranium concentration along length of hair (mg·kg ⁻¹ per length interval), ICPMS	Infant with CBD	Mann-Whitely U-Wilcox non-parametric test	The Fallujah cohort (cases) were found to have significantly higher levels of uranium in scalp hair samples (0.16 U mg·kg ⁻¹ ± 0.11 SD) than the control population (historical control) in Southern Israel (0.062 U mg·kg ⁻¹) (p=0.016); Uranium concentrations in long-hair samples from the Fallujah cases (0.26 U mg·kg ⁻¹ ± 0.09 SD) were found to be more than 2 SD from the mean for control population (historical control) in Northern Sweden (0.057 U mg·kg ⁻¹ ± 0.065 SD), uranium content does not fall along the length of hair in Fallujah cases as compared to Swedish controls, indicating higher exposure among the Fallujah population in the past compared to present

Alborz (2013)	Cross-sectional	2010	Residents of Basrah governorate	Households = 6032 Children = 10,714	Place of residence - Self-reported exposure to “warfare contamination” (Exposed = Yes, Unexposed = No)	Child with CBD	Chi-squared	A significantly higher proportion of children with birth defects in Basrah (105 out of 383) were found to be living in households that reported exposure to “warfare contamination” than children without birth defects (1349 out of 9547) ($p < 0.001$)
Al-Sabbak et al. (2012) (Study 1)	Ecological: Time trend	1994-2011	Births in Al-Basrah city	Population based	Time period (Unexposed = 1994, Exposed = 2003-2011)	CBD incidence rate (number of infants born with CBD per 1000 live births in population)	No statistical test	The CBD incidence rate in Al-Basrah increased 17-fold between 1994 (1.37 CBD per 1000 live births) and 2003 (23 CBD/1000 live births)
Al-Sabbak et al. (2012) (Study 2)	Case-control	May – Aug. 2010	Parents who gave birth to children with CBD at Fallujah General Hospital	Cases = 103 Controls = 9	Uranium concentration in hair ($\mu\text{g}\cdot\text{kg}^{-1}$), ICPMS	Diagnosis of infant at time of delivery at Fallujah General Hospital (Cases = stillbirths or infants with CBD, Controls = healthy live births)	Independent sample t-test	Uranium concentrations in hair from cases (parents of children with birth defects) and was higher than in controls (parents of healthy children), but the difference was not statistically significant ($p > 0.05$)
Al-Sadoon et al. (1999)	Ecological: Time trend	1990-1998	Residents of Basrah	Population based	Time period - Year of congenital anomaly registration (Unexposed = 1990, Exposed = 1991-1998, later years are equated with greater exposure)	CBD incidence	SND test for difference in proportions (z-test)	A significant increase in CBD incidence in Basrah was found between the periods 1991-1994 (2.5 CBDs/1000 live births) and 1995-1998 (4.57 CBDs/1000 live births) (SND=5.37, $p < 0.01$)
Al-Sahlanei et al. (2016)	Case-control	N/R*	Infants born in Baghdad, Dhi-Qar and Basrah.	Participants = 47 mother-neonate pairs	Uranium concentration in maternal and umbilical cord blood samples (ppb), CR-39 fission track detector	Diagnosis at time of delivery in Baghdad (Hospital of Al-Yarmuk and Hospital of Al-Alwiyah), Basrah (Hospital of Gezwan) and Dih-Qar (Hospital of	Independent sample t-tests	Mean uranium concentrations in the maternal and umbilical cord blood samples of deformed infants ($2.43 \text{ ppb} \pm 0.89 \text{ SD}$, and $1.99 \text{ ppb} \pm 0.78 \text{ SD}$, respectively) were found to be significantly higher than those samples from normal infants ($1.26 \text{ ppb} \pm 0.51 \text{ SD}$, and 0.97

						Al-Shatrah and Hospital of Al-Nasriah), (Cases = infants born dead and deformed, Controls = infants born normal and alive)		ppb \pm 0.38 SD, respectively) (p<0.05), samples from Basrah also had significantly higher uranium concentrations than the other two regions (Baghdad and Dhi-Qar)
Neamah & Tawfiq (2015)	Cross-sectional	Jan. 01 – May 31, 2011	Residents of Fallujah	N/R	Place of residence (Exposed = Fallujah, Unexposed = Baghdad)	CBD incidence (number of infants born with CBD per 1000 live births) recorded during a five month period at Fallujah General Hospital (Fallujah) and Yarmouk Teaching Hospital (Baghdad)	Autoregressive model	The coefficient values were found to be higher for Basrah (exposed region) than for Baghdad (unexposed region)
Savabieasfahani et al. (2020)	Case-control	Summer and Fall of 2016	Nasriyah (Bint Al-Huda Maternity Hospital)	Cases = 19 Controls = 10	Uranium and thorium concentration in hair (ICPMS)	Congenital birth defects	ANOVA	The mean concentrations of uranium and thorium in hair samples from cases (43.51 \pm 29.14 and 6.09 \pm 3.22, respectively) and were higher than those from controls, but the differences were not statistically significant
Savabieasfahani et al. (2016)	Case-control	April 2013	Children born with CBDs in Basrah city	Cases = 3 Controls = 6	Uranium and thorium concentrations (ppm) in deciduous teeth, LA-ICP-MS elemental bioimaging	Child with CBD	No statistical test	Uranium and thorium were not detected in any of the samples (detection limit of LA-ICP-MS method was in the ppb range)

Note: CBD = Congenital birth defects, ICP-MS = inductively coupled plasma mass spectrometry, N/R = Not reported; *Paper suggests that the study was conducted after 1991

Table S6. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and **birth-related outcomes** (excluding congenital birth defects) among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Al-Sahlane et al. (2017)	Cross-sectional	N/R*	Infants born at maternity hospitals in Baghdad	Participants = 50 mother-neonate pairs	Uranium concentration in maternal and umbilical cord blood samples (ppm), CR-39 fission track detector	Infant anthropometric measurements (birth weight, body length, head circumference), determined at time of delivery	Adjusted regression model	Uranium concentrations in maternal blood samples ($0.95 \text{ ppm} \pm 0.62 \text{ SD}$) and umbilical cord blood samples ($0.68 \text{ ppm} \pm 0.39 \text{ SD}$) were found to be negatively, significantly correlated with the anthropometric measurements (infant birth weight, body length, and head circumference) ($p < 0.05$), except for infant umbilical cord blood uranium concentrations and body length (correlation was negative but not significant, $p > 0.05$)
Busby et al. (2010) (Study 1)	Cross-sectional	Jan. 20 - Feb. 20, 2010	Residents of Fallujah	Households = 711 Children = 2,132	Time period - Year of birth (Exposed = births after 2005, Unexposed = births prior to 2005)	Birth-sex ratio (ratio of male births to 1000 female births), as reported by subjects in household survey	A statistical test was reportedly used, but not described	Birth-sex ratio decreased to 0.86 for children born between 2006-2010, compared to 1.182 for children born between 2001-2005, 1.109 for children born between 1996-2000, and 1.010 for children born between 1991-1995; the birth-sex ratio for children born between 2006-2010 was found to differ significantly from the expected ratio
Busby et al. (2010) (Study 2)	Case-control	Jan. 20 - Feb. 20, 2010	Residents of Fallujah	Households = 711	Place of residence (Exposed = Fallujah, Unexposed = Egypt, Jordan, and Kuwait)	Infant mortality rate (IMR)	Z-test	IMR in Fallujah between 2006-2010 was four times higher than IMR in Egypt and Jordan ($p < 0.00001$), and nine times higher than the IMR in Kuwait

Note: IMR = Infant mortality rate, N/R = Not reported; *Paper suggests that the study was conducted after 1991

Table S7. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and cancer among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Al-Hamzawi et al. (2015)	Case-control	N/R*	Residents of Southern Iraqi governorates (Basrah, Muthanna, and Dhi-Qar)	Cases = 24 Controls = 12	Uranium concentration in tissue ($\mu\text{g}\cdot\text{kg}^{-1}$), CR-39 fission track detector	Cancer (Cases = kidney, breast, stomach, and uterus cancer tissues, Controls = kidney, breast, stomach, and uterus tissues from healthy individuals)	Independent sample t-test	Significant differences in mean uranium concentrations were found between tissues from cancer patients compared to healthy controls for all cancer types: Kidney ($p < 0.001$) Cancer: $6.51 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.20 \text{ SD}$ Normal: $4.11 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.38 \text{ SD}$ Breast ($p < 0.01$) Cancer: $5.04 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.27 \text{ SD}$ Normal: $2.96 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.16 \text{ SD}$, Stomach ($p < 0.01$) Cancer: $5.22 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.22 \text{ SD}$ Normal: $3.11 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.29 \text{ SD}$ Uterus ($p < 0.01$) Cancer: $4.61 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.32 \text{ SD}$ Normal: $2.28 \mu\text{g}\cdot\text{kg}^{-1} \pm 0.51 \text{ SD}$
Al-Hamzawi et al. (2014)	Case-control	N/R*	Residents of Southern Iraqi governorates (Basrah, Muthanna, and Dhi-Qar)	Cases = 30 Controls = 30	Uranium concentrations in blood samples (ppb), CR-39 fission track detector	Leukemia (Cases = leukemia patients in selected hospitals, Controls = healthy volunteers residing in the same areas as hospitals)	Independent sample t-test	Uranium concentrations in blood samples from the leukemia patients ($2.87 \text{ ppb} \pm 0.11 \text{ SD}$) were found to be significantly higher than those from the healthy group ($1.43 \text{ ppb} \pm 0.07 \text{ SD}$) ($p < 0.001$), and uranium concentrations from cases and controls from Basrah were higher than concentrations in blood samples from the other governorates ($p < 0.05$)
Al-Hashimi & Wang (2013)	Ecological: Time trend	1980-2010	Residents of Ninawa Province	Population based	Time period (Unexposed = 1980-1990, Exposed = 1991-	Cancer incidence rate ratio (IRR)	Poisson regression analysis	IRR for most cancer types in Ninawa significantly decreased in the second (1991-2000) and third period (2001-2010)

					2000 and 2001-2010)			compared to the first period (1980-1990) ($p < 0.01$), leukemia increased in the third period (IRR: 0.2152, CI: 0.1976-0.2346) compared to the second period (IRR: 0.1731, CI: 0.1505-0.1990), but not to the first period (IRR: 0.2964, CI: 0.2433-0.3611)
Al-Jobori (2013)	Case-control	N/R*	Cancer patients residing in the South of Iraq	Cases = 9 Controls = 3	Uranium concentration in tissues (CR-39 fission track detector)	Cancer (Cases = samples from kidney, bone, breast, lung and liver cancer patients, Controls = samples from kidney, bone, and breast cancer patients)	No statistical test	Uranium was not detected in tissue samples from any of the non-cancerous controls
Al-Rudainy et al. (2011)	Ecological: Time trend	2004-2009	Residents of Basrah Governorate	Population based	Time period (Least exposed = 2004, Most exposed = 2009)	Incidence of childhood Leukemia (0-14 years old)	Standard linear regression, test for trend using parameter estimates of regression model	Incidence of childhood leukemia did not change over the 6 year study period. Leukaemia rates decreased by 0.123 per 100,000 between 2004-2009, but the test for trend was not significant ($p=0.81$)
Al-Rudainy et al. (2009a)	Ecological: Geographic comparison	2006	Residents of Basrah Governorate	Population based	Place of residence - Locations of DU contaminated sites in Basrah governorate were compiled through a literature and meta-geographic-analysis	Cancer incidence rate by district	Spearman correlation analysis	No statistical correlation was found between level of DU contamination and cancer incidence rate by district ($r = -0.01$, $p = 0.98$)
Al-Rudainy et al. (2009b)	Ecological: Time trend	2003-2007	Residents of Basrah Governorate	Population based	Time period (Least exposed = 2003, Most exposed = 2007)	Incidence of childhood Leukemia (1-14 years old)	No statistical test	Over the 5 years study period, no increase in childhood leukemia incidence was observed.

Busby et al. (2010) (Study 3)	Case-control	Jan. 20 - Feb. 20, 2010	Residents of Fallujah	Households = 711 Residents = 4,843	Place of residence (Exposed = Fallujah, Unexposed = Egypt, Jordan, and Kuwait)	Cancer incidence	Z-test	Relative Risk (RR) for cancer incidence in Fallujah between 2005-2010 compared to Egypt was 4.22 (CI: 2.8 - 6.6, $p < 0.00000001$)
Hagopian et al. (2010)	Ecological: Time trend	1993-2007	Children (0-14 years of age) residing in Basrah Governorate	Population based	Time period - Year of leukemia registration (recent years equates with greater weaponized uranium exposure, earlier years with less exposure)	Leukemia incidence among children aged 0-14 years, over three year periods	Standard linear regression	A significant ($p=0.03$) trend of increasing incidence of childhood leukemia in Basrah between 1993-2007 was found; incidence more than doubled over the study period (ratio of 2005-2007 incidence to 1993-1995 incidence=2.7; CI=1.437, 5.124)
Hassan et al. (2019)	Case-control	Oct.-Dec. 2017	Cancer patients in Karbala Governorate	Cases = 10 Controls = 2	Uranium concentration in blood samples (ppm), LR-115 type II SSNTD	Cancer	No statistical test	The mean uranium concentration in blood samples from cancer patients (1.4 ppm) was higher than that for healthy controls (0.1 ppm), but no statistical test was performed.
Hassan and Hamadi (2005)	Ecological: Time trend	1997-2002	Residents of Basrah Governorate	Population based	Time period - Year of death or diagnosis (max. exposure = 2002, min. exposure = 1997)	Cancer incidence rate (IR) and mortality rate (MR)	No statistical test reported in study, but a linear regression performed by authors of this SR using reported IR showed a positive trend and moderate effect size (r -squared = 0.58)	No significant increases in cancer IR or MR were observed in Basrah over the study period.
Qaddoori & Shafik (2018)	Case-control	N/R*	Bladder cancer patients in Baghdad Governorate	Cases = 60 Controls = 30	Uranium concentration in urine ($\mu\text{g/L}$) using CR-39 fission track detector.	Bladder cancer	No statistical test	Bladder cancer patients were found to have a higher average concentration of uranium in urine (1.79004 $\mu\text{g/L}$) compared to controls (0.89308 $\mu\text{g/L}$), but the difference was not tested statistically.

Salman (2008)	Ecological: Time trend	1989-2004	Residents of Diyala Governorate	Population based	Time period - Year of cancer diagnosis (Exposed = 2004, Unexposed = 1989)	Number of diagnosed cancer cases per year (Baquba General Hospital, Primary care center of Baquba and medical centers for cancer treatment in Baghdad)	No statistical test	The number of lung cancer cases recorded was higher in 2004 (105 cases) than in 1989 (26 cases), as well as for breast cancer (85 and 17 cases, respectively), and leukemia (92 and 22 cases, respectively) – note: cancer case counts per year do not account for population growth
Shafik (2014)	Case-control	N/R*	Female breast cancer patients in Baghdad	Cases = 41 Controls = 5	Uranium concentrations in 24-hour urine samples ($\mu\text{g}\cdot\text{L}^{-1}$), KPA-11	Breast cancer (Cases = women with breast cancer living in Baghdad, Controls = healthy women living in Baghdad)	No statistical test	The mean concentration of uranium in urine samples was higher among cases (breast cancer patients, $1.6 \mu\text{g}\cdot\text{L}^{-1} \pm 0.027 \text{ SD}$) than controls (healthy women, $1.03 \mu\text{g}\cdot\text{L}^{-1} \pm 0.0202 \text{ SD}$)
Showard & Aswood (2019)	Case-control	N/R*	Patients at Morgan Hospital in Babylon Governorate	Cases = 24 Controls = 6	Concentration of radon-emitted alpha particles in blood samples (Bq/m^3), CR-39 fission track detector	Leukemia	No statistical test	The concentration of alpha particles in blood samples for leukemia patients was higher than for controls (7.79 Bq/m^3 and 4.39 Bq/m^3 , respectively), but the difference was not tested statistically.

Note: SSNTD = Solid state nuclear track detector, N/R = Not reported; *Paper suggests that the study was conducted after 2003

Table S8. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and immune system function among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Abdul-Wahid (2009)	Case-control	June 2007	Residents of a district in northern Al-Basrah city	Cases = 50 Controls = 50	Place of residence (Exposed = Basrah, Unexposed = Baghdad)	Immune system function (lymphocyte phenotyping: % of cells as lymphocytes in blood sample)	T-test	Cases were found to have lower levels of selected lymphocytes (CD3, CD4, CD8, CD19, and CD56) compared to controls (p-values not reported)
Humaidi & Khalaf (2011)	Case-control	Aug. 2005 – Aug. 2009	Bullet wounded Iraqi's in Ramadi	Cases = 196 Controls = 19	Bullet type (Exposed = shot by US or coalition forces, unexposed = shot by other source)	W.B.C. count, hemoglobin concentration, erythrocyte sedimentation ratio, total serum Bilirubin, alkaline phosphates enzyme concentration, serum transferees enzymes concentration, serum Creatinine concentration, blood urea concentration, mitotic index.	ANOVA	There were significant differences (p<0.05) between cases and controls for all outcomes measured
Mryoush & Salim (2015)	Cross-sectional	N/R**	Residents of Baghdad	Participants = 50	Place of residence - Uranium concentration in soil samples (ppm) from five neighborhoods in Baghdad (North -	Mitotic index (MI) analysis (number of cells undergoing mitosis/1000 cells in blood sample)	No statistical test	The North of Baghdad had the highest mean uranium concentration in soil samples (12.90 ppm ± 0.7 SD) and the West had the lowest mean (0.60 ppm ± 0.21 SD), and the mean Mitotic Index in blood samples from the North (2.3 ± 0.059 SD)

					Al-Taji, East - Diyala Bridge, South - Al- Mhmodya, West - Abu Ghraib, Central - Bab-Al- Sharqee)			was higher than the mean MI in samples from the West (0.20 ± 0.3 SD), suggesting a negative correlation
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Note: W.B.C. = White blood cell, N/R = Not reported; **Paper suggests that the study was conducted after 1991

Table S9. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and **kidney failure** among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Abed et al (2019, 2020)	Case-control	N/R*	Al-Muthanna Governorate	Cases=21 Controls=5	Uranium concentration in blood (ppm) and urine($\mu\text{g/L}$), CR-39 nuclear track detector	Kidney failure	No statistical test	The mean uranium concentration in blood samples from kidney failure patients (0.243 ppm) was higher than that for healthy controls (0.137 ppm), but no statistical test was performed. The mean uranium concentration in urine samples from kidney failure patients (1.90 $\mu\text{g/L}$) was higher than that for healthy controls (1.16 $\mu\text{g/L}$), but no statistical test was performed.

Note: N/R= Not reported; * Paper suggests that the study was conducted after 2003

Table S10. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and **BCL-2 oncogene expression** among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Mohammad (2016)	Case-control	2007-2009	Breast cancer patients in Baghdad	Cases = 50 Controls = 30	Place of residence (Exposed = Iraq, Unexposed = Italy)	Bcl-2 oncogene expression and intensity in breast cancer tissue samples	Chi-squared (for Bcl-2 expression) No statistical test (for Bcl-2 intensity)	Bcl-2 expression in Iraqi breast cancer tissue samples was found to be significantly higher ($p = 0.037$) than in Italian samples, and among individuals for which Bcl-2 was positively expressed, Iraqi participants had higher intensities than Italian participants

Table S11. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and **PTEN gene expression** among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Jumaah et al. (2019)	Case-control	Oct. 2006 - Oct. 2007	Female patients at hospitals in central and southern Iraq	Cases (exposed) = 21 Controls (unexposed) = 22	Place of residence (conflict zone vs. peaceful area)	PTEN gene expression	Student's T-test	The PTEN gene expression mean fold change was greater among the exposed group (0.139 ± 0.185) than the unexposed group (0.0031 ± 0.0029), but the difference in means was not statistically significant when tumor grade and cancer stage were controlled for ($p=0.286$ and $p=0.98$, respectively).

Note: N/R= Not reported; * Paper suggests that the study was conducted after 2003

Table S12. Study characteristics of human observational studies measuring association between exposure to weaponized uranium and **mixed adverse health outcomes** among the Iraqi population

Study ID	Study Design	Study Timeframe	Population/ Location	Sample size	Exposure	Outcome(s)	Statistical test(s)	Results
Al-Hamadany et al. (2012) (Study 1)	Case-control	2009-2010	Residents of Baghdad	Cases = 74 Controls = 14	Uranium concentration in blood samples (ppm), CR-39 fission track detector	Illness (Cases = patients with cancer and mothers of children with CBD, Controls = healthy adults)	T-test	Results comparing predefined cases and controls were not reported; Rather, mean uranium concentrations in blood samples are compared between a portion of the control samples (only those healthy individuals living in uncontaminated areas) (0.11ppm ± 0.009 SE), and the cancer patient samples plus the other controls (0.21 ppm ± 0.01 SE) and the difference was statistically significant (p<0.05)
Al-Hamadany et al. (2012) (Study 2)	Case-control	2009-2010	Residents of Baghdad	Cases = 74 Controls = 14	Place of residence, health status, or occupation (Exposed = cancer patients, mothers of children with CBDs, employees of the Institute and Hospital of Radiotherapy and Nuclear Medicine, or individuals	Total and Differential W.B.C. Count, Hemoglobin Concentration, neutrophils phagocytic activity, IFN-γ concentrations, IL-2 concentrations	Independent sample t-tests	WBC counts were significantly higher among groups defined as “exposed” and hemoglobin concentrations were significantly lower compared to the group defined as unexposed (p<0.05)

					residing in areas of Baghdad identified by UNEP as contaminated, Unexposed = healthy individuals residing in areas reported to be free of weaponized uranium)			
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Table S13. Risk of bias assessment for Abed et al (2019, 2020) (Reference 27)

Domain	Rating	Justification
Recruitment	Probably low risk	The recruitment methods in this study were not fully described, but it is suggested that participants were selected from the same area during the same time period.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Low risk	The method for uranium exposure measurement (CR-39 fission track detector) is robust.
Outcome Assessment	Probably low risk	The authors do not sufficiently describe how kidney failure patients were diagnosed, but the use of the term 'patients' suggests that they were diagnosed by medical professionals.
Confounding	High risk	The study collected data on (but did not control for) age and sex of participants. No other confounders were controlled for.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, "The Navigation Guide instructions for making risk of bias determinations"

Table S14. Risk of bias assessment for Abdul-Wahid (2009) (Reference 1)

Domain	Rating	Justification
Recruitment	Probably high risk	Study lacks complete description of recruitment criteria, but it is suggested that participants from the unexposed population were selected to be predisposed towards normal (“healthy”) immune system function, while the same criteria was not applied to the selection of participants from the exposed population.
Blinding	Probably high risk	Study does not report blinding of key personnel (e.g. personnel counting the number of labeled cells).
Exposure Assessment	High risk	The city of Baghdad does not represent an unexposed geographic location with certainty, as heavy fighting took place in the city during the 2003 invasion by US and coalition forces.
Outcome Assessment	Probably low risk	Outcomes were assessed and defined consistently across all study participants, using a valid and reliable measures (biomarkers) – no Quality assurance/Quality control.
Confounding	Probably high risk	The study controlled for age and sex, but it is unclear whether is accounted to for other important confounders including tobacco use and obesity.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Low risk	The study is free of suggestion of selective outcome reporting. All of the study’s specified outcomes were adequately reported.
Other Bias	High risk	The study reportedly performed t-test to measure association between cases and controls, but the p-values for the test were reported inaccurately.
Conflict of Interest	Low risk	The authors report no conflict of interest, and associated funds and persons appear to be from government and/or academia only.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S15. Risk of bias assessment for Alaani et al (2012) (Reference 14)

Domain	Rating	Justification
Recruitment	High risk	Recruitment strategies for the control populations are not described.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably high risk	While Egypt (Giza) and Jordan represent unexposed populations, depleted uranium has previously been detected in Kuwait.
Outcome Assessment	Probably low risk	Diagnoses of congenital anomalies were made a professional pediatrician (consistent and valid) – no Quality assurance/Quality control.
Confounding	High risk	The study used a questionnaire to collect data on most important confounders among study participants. However, no confounding variables were accounted for between study groups.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Low risk	The study is free of suggestion of selective outcome reporting. All of the study's specified outcomes were adequately reported.
Other Bias	Probably low risk	The method for assessing the incidence of congenital birth defects per 1000 live births in Fallujah was not precise (the denominator was estimated), but the explanation provided in the report suggests that the method was reasonably accurate.
Conflict of Interest	High risk	Authors report no conflict of interest, but funding for the study was partially provided by Swedish non-profit International Foundation for Research on Radiation Risk (IFRRR), which has a stated agenda of disputing the

		International Commission on Radiological Protection (ICRP) radiation risk model.
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Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S16. Risk of bias assessment for Alaani et al (2011) (Reference 13)

Domain	Rating	Justification
Recruitment	High risk	Cases and controls were recruited from different populations (Iraq, Israel, and Sweden). Descriptions of the recruitment strategies for the Israeli population (historical control) and Swedish population (historical control) are not reported, nor is the recruitment criteria for Fallujah participants fully reported.
Blinding	Low risk	Hair samples for uranium concentration analysis were re-coded to ensure blinding of key personnel.
Exposure Assessment	Low risk	The method of uranium concentration measurement in hair samples using Inductively coupled plasma mass spectrometry (ICMPS) is robust.
Outcome Assessment	Probably high risk	Cases were diagnosed by a medical doctor at Fallujah General Hospital, however it is not known if or how outcomes were assessed among the participants in the comparator group (Southern Israel, historical control). It is not known if or how outcomes were assessed among the participants in the Swedish (historical control) comparator group.
Confounding	High risk	The study did not account for many important confounders including consanguinity, obesity, maternal folate deficiency or maternal education.
Incomplete Outcome Data	High risk	Birth outcomes were not reported for either historical control group (Israeli and Swedish).
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	High risk	Authors report no conflict of interest, but funding for the study was partially provided by Swedish non-profit

		International Foundation for Research on Radiation Risk (IFRRR), which has a stated agenda of disputing the International Commission on Radiological Protection (ICRP) radiation risk model.
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Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S17. Risk of bias assessment for Alborz (2013) (Reference 15)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	High risk	Self-reported exposure to war contamination is not a robust method of measuring uranium exposure.
Outcome Assessment	Probably low risk	Outcomes were assessed and defined consistently across all study participants, using a method (questionnaires) that were valid and reliable for the outcome of interest (birth defects) – no Quality assurance/Quality control.
Confounding	High risk	Although the study accounted to other environmental exposures, it did not account for any other important confounders.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S18. Risk of bias assessment for Al-Hamadany et al (2012) (Study 1) (Reference 2)

Domain	Rating	Justification
Recruitment	Probably low risk	Study lacks a complete description of recruitment methods.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Low risk	The method for uranium exposure measurement (CR-39 fission track detector) is robust.
Outcome Assessment	Probably high risk	The study does not clearly define the outcome of interest or report how the outcomes were assessed.
Confounding	High risk	Study did not control for any confounders.
Incomplete Outcome Data	Low risk	Study appears to be free of missing outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	High risk	The study did not report uranium concentrations for all six study groups. The decision to lump occupationally exposed individuals, and healthy individuals living in neighborhoods suspected to be contaminated with depleted uranium, into the same group as cancer patients and mothers who gave birth to children with congenital birth defects (cases) for comparison to the health, unexposed group, introduces a serious methodological flaw into the study.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S19. Risk of bias assessment for Al-Hamadany et al (2012) (Study 2) (Reference 2)

Domain	Rating	Justification
Recruitment	Probably low risk	Study lacks a complete description of recruitment methods.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	High risk	The study defined uranium “exposed” cases as cancer patients, mothers of children with congenital birth defects, employees of the Institute and Hospital of Radiotherapy and Nuclear Medicine, or individuals residing in areas of Baghdad identified by the United Nations Environmental Program (UNEP) as contaminated. This is a highly problematic definition of “exposed”.
Outcome Assessment	Probably low risk	The methods used for outcome assessments (direct assessment of biomarkers) were valid and robust – no Quality assurance/Quality control .
Confounding	High risk	Study did not control for any confounders.
Incomplete Outcome Data	Low risk	Study appears to be free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears to be free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S20. Risk of bias assessment for Al-Hamzawi et al (2015) (Reference 4)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Low risk	Tissue samples for uranium concentration analysis were re-coded to ensure blinding of key personnel.
Exposure Assessment	Low risk	The methods for uranium concentration measurement in blood samples (CR-39 fission track detector) are robust.
Outcome Assessment	Probably low risk	All samples were collected from a histopathology clinic in Southern Iraq, which presumably used valid and reliable methods to record or determine if samples came from subjects diagnosed with cancer, or healthy subjects – No Quality assurance/Quality control .
Confounding	High risk	Data on age, sex, and tobacco use of participants was collected and reported. However, the confounders were not accounted or controlled for in analysis.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	High risk	Neither the time frame during which tissue samples of cancer patients (cases) were collected by the histopathology clinic, nor the year that cancer was diagnosed for study subjects are reported. Cancerous tissue samples collected before 1991 or 2003, or collected from patients who were diagnosed before 1991 or 2003, cannot inform the question of whether uranium exposure (independent variable) is association with cancer (dependent variable). Samples from patients with inherited-type cancers also cannot inform the question of association. Some cancer treatment drugs

		can affect kidney function, which could hypothetically reduce the excretion rate of uranium, leading to higher concentrations of uranium in tissues among patients receiving treatment than in healthy volunteers, although the levels of environmental exposure may be the same. Likewise, cancer of the kidney can impair kidney function, leading to a reduction in the rate of uranium excretion and an accumulation of uranium in kidney tissues.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S21. Risk of bias assessment for Al-Hamzawi et al (2014) (Reference 3)

Domain	Rating	Justification
Recruitment	Probably low risk	Study participants were not all recruited from the same population, but proportions of participants from each population in each study group are uniform. Study lacks a complete description of recruitment methods, but otherwise no reason to suspect there were substantial differences between comparison groups other than uranium exposure.
Blinding	Low risk	Blinding methods are not reported, but the authors of this SR judge that the neither the outcome and the outcome measurement, nor exposure and exposure measurement are likely to be influenced by lack of blinding.
Exposure Assessment	Low risk	The methods for uranium concentration measurement in blood samples (CR-39 fission track detector) are robust.
Outcome Assessment	High risk	The outcome of interest (Leukemia) was not assessed consistently across all study participants.
Confounding	High risk	The study accounted for age and sex of participants, but it did not account for any other important confounders.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	High risk	Neither the year that blood samples of leukemia patients were collected by hospitals, nor the year that cancer was diagnosed for the leukemia patients are reported. Leukemia blood samples collected before 1991 or 2003, or collected from patients who were diagnosed before 1991 or 2003, cannot inform the question of whether uranium exposure (independent variable) is association

		with leukemia (dependent variable). Samples from patients with inherited-type leukemia also cannot inform the question of association. Some leukemia treatment drugs can affect kidney function, which could hypothetically reduce the excretion rate of uranium, leading to higher concentrations of uranium in blood among patients receiving treatment than in healthy volunteers, although the levels of environmental exposure may be the same.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S22. Risk of bias assessment for Al-Hashimi & Wang (2013) (Reference 5)

Domain	Rating	Justification
Recruitment	Probably low risk	Data for this study was obtained from the Directorate of Health in Ninawa. The methods by which the Directorate collected the data between years was not reported, but there is no suggestion that methods of data collection differed between years.
Blinding	Not applicable	As a purely statistical analysis, blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Low risk	Given the study design (ecological: time trend), the method of exposure measurement in this study (year) is robust.
Outcome Assessment	Probably low risk	Data was collected from the Directorate of Health of the Ninawa province in Iraq.
Confounding	High risk	While the study controlled for age and sex in its analysis, it did not control for tobacco use or other environmental exposures.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Low risk	The study is free of suggestion of selective outcome reporting. All of the study's specified outcomes were adequately reported.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Low risk	The authors report no conflict of interest, and associated funds and persons appear to be from government and/or academia only.

Note: For more information, please see Supplemental Material, "The Navigation Guide instructions for making risk of bias determinations"

Table S23. Risk of bias assessment for Al-Jobori (2013) (Reference 6)

Domain	Rating	Justification
Recruitment	High risk	Participants were not recruitments not recruited from the same population. The timeframe during which participants were recruited is not reported.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Probably high risk	While the method used for uranium concentration measurement in tissues samples (CR-39 fission track detector) is robust, the finding that control samples contained no detectable levels of uranium is highly questionable.
Outcome Assessment	Probably low risk	All subjects (cases and controls) were recruited from hospitals in Iraq where the health outcome of interest (cancer) was diagnosed by a medical doctor.
Confounding	High risk	Study did not account for any confounders.
Incomplete Outcome Data	Low risk	Study appears free from missing outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	High risk	The study did not clearly state whether the specimens were collected from affected organs. The sample size was small (controls, n=3).
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S24. Risk of bias assessment for Al-Rudainy et al (2011) (Reference 16)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably high risk	The time frame for this study was 2004-2009, and thus no data for an unexposed population (prior to 1991 or 2003) was used in this study. The assumption that later year of diagnosis (more recent) equates with higher uranium exposure is questionable.
Outcome Assessment	Low risk	Health outcome data was collected from the Pediatric Oncology Ward in Basrah Maternity & Children's Hospital and the Basrah Health Authorities Statistical Office.
Confounding	High risk	While the study controlled for age and sex in its analysis, it did not control for tobacco use or other environmental exposures.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, "The Navigation Guide instructions for making risk of bias determinations"

Table S25. Risk of bias assessment for Al-Rudainy et al (2009a) (Reference 7)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably low risk	Given the study design (ecological: geographic comparison), the method of exposure measurement in this study (meta-synthesis of depleted uranium impacted sites) is robust.
Outcome Assessment	Low risk	Health and population data was compiled from a valid and reliable source.
Confounding	High risk	While the study controlled for age and sex in its analysis, it did not control for tobacco use or other environmental exposures.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S26. Risk of bias assessment for Al-Rudainy et al (2009b) (Reference 8)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably high risk	The time frame for this study was 2007-2009, and thus no data for an unexposed population (prior to 1991 or 2003) was used in this study. The assumption that later year of diagnosis (more recent) equates with higher uranium exposure is questionable.
Outcome Assessment	Low risk	Health outcome data was collected from the Pediatric Oncology Ward in Basrah Maternity & Children's Hospital and the Basrah Health Authorities Statistical Office.
Confounding	High risk	Study did not account for tobacco use among participants or population, nor other environmental exposures.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Low risk	Authors declared no conflict of interest. Associated funds and persons appear to be from academia only and free of financial interests in study results.

Note: For more information, please see Supplemental Material, "The Navigation Guide instructions for making risk of bias determinations"

Table S27. Risk of bias assessment for Al-Sabbak et al (2012) (Study 1) (Reference 9)

Domain	Rating	Justification
Recruitment	Low risk	Recruitment criteria were consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably high risk	The control period (1994) does not represent an unexposed period, as depleted uranium weapons were used in Basra by US forces in 1991.
Outcome Assessment	High risk	Data was obtained from the Al-Basrah Maternity Hospital (a reliable source). However, health outcomes among the control population were only assessed over a 1 year period, compared to the exposed population for which birth defects incidence was calculated over a 9 year period. The impact of economic sanctions in Iraq on health care infrastructure during that period (1994) may have impacted cancer surveillance and registration.
Confounding	High risk	No confounding variables in the study population were measured or accounted for.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Low risk	The study is free of suggestion of selective outcome reporting. All of the study's specified outcomes were adequately reported.
Other Bias	High risk	Only a fraction of the total births in Al-Basrah take place in a hospital setting, or at Al-Basrah Maternity hospital specifically, which could introduce a form of selection bias into the study design.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial

		interests in study results. However, no claim denying conflicts of interest was made.
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Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S28. Risk of bias assessment for Al-Sabbak et al (2012) (Study 2) (Reference 9)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Probably high risk	The study does not discuss blinding of key personnel (e.g. personnel conducting ICPMS analysis), and it is possible that lack of blinding could introduce bias.
Exposure Assessment	Probably low risk	The method of uranium concentration measurement in hair samples using Inductively coupled plasma mass spectrometry (ICPMS) is robust, as long as hair samples were collected from the scalp (not specified in study).
Outcome Assessment	Low risk	Diagnosis of birth defects by a medical doctor at the Fallujah General Hospital at the time of delivery is a valid and robust methods for outcome assessment.
Confounding	High risk	A questionnaire was reportedly used to collect data on many important confounders (including consanguinity and maternal health status) and the authors also tested hair samples for other environmental teratogens including lead and mercury. However, these confounding variables to do appear to have been accounted for in the statistical analysis comparing uranium concentration in hair samples between cases and controls.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Low risk	The study is free of suggestion of selective outcome reporting. All of the study's specified outcomes were adequately reported.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial

		interests in study results. However, no claim denying conflicts of interest was made.
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Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S29. Risk of bias assessment for Al-Sadoon et al (1999) (Reference 10)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Low risk	Given the study design (ecological: time trend), the method of exposure measurement in this study (year) is robust.
Outcome Assessment	Low risk	Data was obtained from the Al-Basrah Maternity Hospital (a reliable source).
Confounding	High risk	No confounding variables in the study population were measured or accounted for.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S30. Risk of bias assessment for Al-Sahlanee et al (2017) (Reference 12)

Domain	Rating	Justification
Recruitment	Probably low risk	Study lacks a complete description of recruitment criteria. Namely, it does not specify if participants were recruited from a single hospital or multiple hospitals in Baghdad, or explicitly define the time period during which participants were recruited.
Blinding	Probably high risk	The study does not discuss blinding of key personnel (e.g. personnel counting CR-39 fission detector tracks), and it is possible that lack of blinding could introduce bias.
Exposure Assessment	Low risk	The methods for measuring uranium concentration measurement in blood samples (CR-39 fission track detector) are robust.
Outcome Assessment	Probably low risk	Details on the methods used to obtain anthropometric of measurements of newborns are lacking, but presumably performed by a nurse at the hospital at the time of birth – No Quality assurance/Quality control.
Confounding	Probably high risk	Data on all important confounders except consanguinity were collected. It is suggested (but not explicitly stated) that they were controlled for in the adjusted regression analysis.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All of the pre-specified outcomes are reported.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S31. Risk of bias assessment for Al-Sahlanee et al (2016) (Reference 11)

Domain	Rating	Justification
Recruitment	High risk	Study lacks a complete description of recruitment criteria. Participants were recruited from different populations, and the number of participants from each population are not reported. The time period during which blood samples were collected from participants is not reported.
Blinding	Probably high risk	Blinding of key personnel (e.g. personnel conducting track density counts) was not reported.
Exposure Assessment	Low risk	The method of uranium concentration measurement in maternal and umbilical cord blood samples (CR-39 fission track detector) is robust.
Outcome Assessment	High risk	Details are lacking on how birth outcomes were defined or diagnosed during this study. It unclear whether cases included still births without diagnosed congenital malformations.
Confounding	High risk	Data on maternal age was collected, but not controlled for in analysis.
Incomplete Outcome Data	Low risk	The study did not have incomplete outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S32. Risk of bias assessment for Busby et al (2010) (Study 1) (Reference 17)

Domain	Rating	Justification
Recruitment	Probably low risk	Recruitment criteria was applied similarly across study groups. The response rate for the household survey was reportedly 60%, and the majority of the non-responses came from a single neighborhood where household residents were suspicious of the surveyors. The study authors provide a reasonable explanation that the non-responses were unlikely to be related to exposures or outcomes.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Low risk	Given the study design (cross-sectional), the method of exposure measurement in this study (year) is robust.
Outcome Assessment	High risk	This study determined the yearly birth-sex ratio in Fallujah by conducting a household cross-sectional survey, and document the age and sex of children in each household. This is a problematic method, because it does not account for deaths, adoptions, or immigration. The study design was also potentially subject to recall bias or overreporting bias. Differential child mortality rates between sexes could also impact outcomes measured.
Confounding	High risk	No confounding variables in the study population were measured or accounted for.
Incomplete Outcome Data	Probably low risk	The study does not report any missing data (but they also do not report that all questionnaires were completed in full).
Selective Reporting	High risk	The study does not report the birth-sex ratio for age cohorts above 19 years. The next age cohort (20-24 years) has a birth-sex ratio even lower than the 0-4 years age cohort (776 compared to 860 males per 1000 females, respectively).

Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Low risk	Authors declare no conflict of interest.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S33. Risk of bias assessment for Busby et al (2010) (Study 2) (Reference 17)

Domain	Rating	Justification
Recruitment	High risk	The recruitment strategies differed between exposed (Fallujah) and unexposed (Egypt) populations.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Low Risk	Egypt represents an unexposed population.
Outcome Assessment	High Risk	The method of data collection among the exposed population (Fallujah) was indirect (cross-sectional survey) potentially subject to recall bias or overreporting bias.
Confounding	High risk	No confounding variables in the study population were measured or accounted for.
Incomplete Outcome Data	Probably low risk	The study does not report any missing data (but they do not report that all questionnaires were completed in full). The study suggests that parents may have underreported cases of birth defects, but the study accounted for that by collecting data on still births (with the reasonable assumption that families effected by stigma surrounding birth defects would report cases as still births or infant mortality).
Selective Reporting	Probably high risk	The questionnaire used in the study aimed to collect data on health status, birth history, and infant mortality among study participants for a ten year period, but only the most recent five year period was reported.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Low risk	Authors declare no conflict of interest.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S34. Risk of bias assessment for Busby et al (2010) (Study 3) (Reference 17)

Domain	Rating	Justification
Recruitment	High risk	The recruitment strategies differed between exposed (Fallujah) and unexposed (Egypt) populations.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Low risk	Egypt represents an unexposed population.
Outcome Assessment	High risk	The method of data collection differed across study groups, and the method of data collection among the exposed population (Fallujah) was potentially subject to recall bias or overreporting bias.
Confounding	High risk	The only confounding variable controlled for in this study was age.
Incomplete Outcome Data	Probably low risk	The study does not report any missing data (but they do not report that all questionnaires were completed in full).
Selective Reporting	Probably high risk	The questionnaire used in the study aimed to collect data on health status, birth history, and cancer history among study participants for a ten year period, but only the most recent five year period was reported.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Low risk	Authors declare no conflict of interest.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S35. Risk of bias assessment for Hagopian et al (2010) (Reference 18)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably high risk	The time frame for this study was 1993-2007, and thus no data for an unexposed population (prior to 1991) was used in this study. The assumption that later year of diagnosis (more recent) equates with higher uranium exposure is questionable.
Outcome Assessment	Low risk	Data was obtained from the Ibn Ghazwan Hospital's leukemia registry.
Confounding	High risk	No confounding variables in the study population were measured or accounted for.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, "The Navigation Guide instructions for making risk of bias determinations"

Table S36. Risk of bias assessment for Hassan et al (2019) (Reference 29)

Domain	Rating	Justification
Recruitment	Probably low risk	The recruitment methods in this study were not fully described, but it is suggested that participants were selected from the same area during the same time period.
Blinding	Probably low risk	It is reported that samples were recoded, which suggests that blinding procedures were implemented.
Exposure Assessment	Low risk	The method for uranium exposure measurement (Solid State Nuclear Track Detector) is robust.
Outcome Assessment	Probably low risk	Patients were reportedly diagnosed at a cancer hospital in Karbala Governorate.
Confounding	High risk	The study did not control for any confounders.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S37. Risk of bias assessment for Hassan et al (2005) (Reference 19)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	Probably high risk	The cancer registry from which data was obtained for this study was created in 1997, and thus no data for an unexposed population was used in this study. The assumption that later year of diagnosis (more recent) equates with higher uranium exposure is questionable.
Outcome Assessment	Low risk	Data was obtained from the Basrah Oncology Center, Basrah Health Office, and Central Statistical Bureau.
Confounding	High risk	Study accounts for sex, but does not account for average age of population between years (increase in cancer rate is expected with an ageing population). No other confounders were accounted for.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S38. Risk of bias assessment for Humaidi & Khalaf (2011) (Reference 20)

Domain	Rating	Justification
Recruitment	Probably low risk	Study lacks a complete description of recruitment methods, but otherwise no reason to suspect there were substantial differences between comparison groups other than uranium exposure (as defined within the context of the study).
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	High risk	Not all bullets used by US or coalition forces contain weaponized uranium.
Outcome Assessment	Probably low risk	The methods used for outcome assessments (direct assessment of biomarkers) were valid and robust – no Quality assurance/Quality control.
Confounding	High risk	Study does not account for age, severity of bullet injury, or location of injury.
Incomplete Outcome Data	Low risk	Study has no suggestion of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears to be free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S39. Risk of bias assessment for Jumaah et al (2019) (Reference 30)

Domain	Rating	Justification
Recruitment	Low risk	Recruitment strategies were the same across study groups.
Blinding	Low risk	Participant data was coded and blinding procedures were implemented.
Exposure Assessment	Probably high risk	Categorization of place of residence as ‘exposed’ or ‘unexposed’ was not verified by environmental or biological monitoring.
Outcome Assessment	Low risk	Study reports that “All cases were examined by two independent pathologists to confirm the diagnosis.”
Confounding	Probably high risk	The study controlled for age and sex of participants, as well as other environmental exposures. It did not control for tobacco use or obesity.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Low risk	The authors declare no funding for this study and no competing interests.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S40. Risk of bias assessment for Mohammad (2016) (Reference 23)

Domain	Rating	Justification
Recruitment	Probably low risk	Study lacks complete description of recruitment criteria for each population, but it is reported that samples were collected from patients in both populations during the same time frame.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Low risk	The Italian control population represents an appropriate (unexposed) control.
Outcome Assessment	Probably high risk	The study reports that the difference in tissue processing time, the difference in tissue quality, and the timing for embedding tissues in paraffin may have impacted outcome measurements, but data is not available for those variables - no Quality assurance/Quality control reported.
Confounding	High risk	Only age and sex were accounted for in this study.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	Low risk	The study appears to be free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S41. Risk of bias assessment for Mryoush & Salim (2015) (Reference 21)

Domain	Rating	Justification
Recruitment	Probably low risk	Study lacks a complete description of recruitment criteria, but otherwise no reason to suspect there were substantial differences between comparison groups other than uranium exposure.
Blinding	Probably high risk	Blinding of key personnel for either exposure or outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Low risk	The method of measuring uranium concentration in soil samples (CR-39 fission track detector) is robust.
Outcome Assessment	Probably low risk	Outcomes were assessed and defined consistently across all study participants, using valid and reliable measures – no Quality assurance/Quality control reported.
Confounding	Probably high risk	The study accounted for age, sex, and tobacco use among study participants. However, it did not account for body fat (obesity) or other environmental exposures.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All of the pre-specified outcomes are reported.
Other Bias	High risk	The authors of the study reported that the use of phosphate fertilizers, or proximity to fertilizer production facilities, could lead to higher concentrations or uranium in soil samples in the study area.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from government and/or academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S42. Risk of bias assessment for Neamah & Tawfiq (2015) (Reference 22)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	High risk	The city of Baghdad does not represent an unexposed geographic location with certainty, as heavy fighting took place in the city during the 2003 invasion by US and coalition forces.
Outcome Assessment	Probably low risk	Not enough information to permit a judgement of low risk of bias, but the article suggests that outcomes were assessed and defined consistently across all study participants, using valid and reliable measures (direct observation by medical professional in prospective cohort study).
Confounding	High risk	The study did not control for any confounders.
Incomplete Outcome Data	Probably low risk	The data in this study is presented in the form of matrices prepared for statistical analysis, from which missing outcome data cannot be interpreted. The authors do not report any missing outcome data in the narrative text.
Selective Reporting	High risk	The study does not report total number of births at each hospital.
Other Bias	Low risk	The study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S43. Risk of bias assessment for Qaddoori & Shafik (2018) (Reference 31)

Domain	Rating	Justification
Recruitment	Probably low risk	The recruitment methods in this study were not fully described, but it is suggested that participants were selected from the same area during the same time period.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Low risk	The method of uranium concentration measures (CR-39 fission track detector) was robust.
Outcome Assessment	Probably low risk	All cancer patients were recruited from hospitals in the Baghdad governorate.
Confounding	High Risk	The study did not control for any confounders.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S44. Risk of bias assessment for Salman (2008) (Reference 24)

Domain	Rating	Justification
Recruitment	Low risk	The strategy for recruiting participants was consistent across study groups.
Blinding	Not applicable	Blinding is not an element of study design capable of introducing risk of bias in the study.
Exposure Assessment	High risk	Study findings may reflect changes in cancer detection (diagnostic abilities), and not true change of cancer incidence in the population.
Outcome Assessment	Low risk	Data on cancer cases was obtained from the Baquba General Hospital.
Confounding	High risk	The study did not control for any confounders.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes are reported.
Other Bias	High risk	Study reports cases, not incidence of cancer.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S45. Risk of bias assessment for Savabieasfahani et al (2020) (Reference 32)

Domain	Rating	Justification
Recruitment	Probably high risk	Recruitment strategies were the same across study groups, but a high non-response rate was reported (39%).
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Low risk	The method of uranium and thorium detection in hair samples Inductively coupled plasma mass spectrometry (ICP-MS) was robust.
Outcome Assessment	Low risk	Study reports that “Determination of congenital anomalies was done by medical doctors at Bint Al-Huda Maternity Hospital.”
Confounding	High risk	The study controlled for age and sex of participants, tobacco use, maternal health status, and maternal age at birth. However, it did not assess or control for maternal folate deficiency, consanguinity, or other environmental exposures.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	High risk	The article states that 43 families were recruited into the study. However, results for only 30 participants are reported (20 cases, 10 controls) were reported.
Other Bias	High risk	The Journal Pre-Proof of this article (published online in August 2019) identified the US military base of interest as Camp Taji. When published, the article was revised and ‘Camp Taji’ was replaced with “Tallil Air Base’ without any explanation. This raises concerns

		about the certainty in identification of the US military base. Additionally, Table 1, reported 20 cases were included in the study, but the manuscript text reports that 19 cases were included (Section 6. Biological Samples).
Conflict of Interest	Low risk	The authors declare no competing interest.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S46. Risk of bias assessment for Savabieasfahani et al (2016) (Reference 25)

Domain	Rating	Justification
Recruitment	High risk	Participants from the control group (children without congenital birth defects) were selected from populations outside of Iraq.
Blinding	Probably high risk	Blinding of key personnel for outcome measurement is not reported, and lack of blinding could introduce bias.
Exposure Assessment	Probably high risk	The method selected for exposure assessment (elemental bioimaging using laser ablation Inductively coupled plasma mass spectrometry) may have a detection limit above the expected range of uranium concentration in human teeth.
Outcome Assessment	Low risk	Outcomes were assessed and defined consistently across all study participants, using valid and reliable measures.
Confounding	High risk	The study did not control for any confounders.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	High risk	Very small sample sizes (Cases, n=3; Controls, n=6)
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S47. Risk of bias assessment for Shafik (2014) (Reference 26)

Domain	Rating	Justification
Recruitment	Low risk	Study lacks a complete description of recruitment methods, but otherwise no reason to suspect there were substantial differences between comparison groups other than uranium exposure.
Blinding	Low risk	Urine samples for uranium concentration analysis were re-coded to ensure blinding of key personnel.
Exposure Assessment	Low risk	The method for monitoring uranium concentration in urine samples (Kinetic Phosphorescence Analyzer, KPA-11) is robust.
Outcome Assessment	Probably low risk	Study lacks a complete description of outcome assessment methods.
Confounding	High risk	Of the important confounders pre-specified in the present systematic review, Shafik (2014) only controlled for sex of participants and tobacco use. The study did not control for age, obesity, or other environmental exposures.
Incomplete Outcome Data	Low risk	Study was free of missing outcome data.
Selective Reporting	Not applicable	In this case-control study, cases were selected on the basis of outcome, thus selective outcome reporting is not capable of introducing risk of bias in the study.
Other Bias	High risk	Very small sample size (Controls, n=5). No statistical test of difference in uranium concentrations was performed.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, “The Navigation Guide instructions for making risk of bias determinations”

Table S48. Risk of bias assessment for Showard & Aswood (2019) (Reference 33)

Domain	Rating	Justification
Recruitment	Probably low risk	The recruitment methods in this study were not fully described, but it is suggested that participants were selected from the same area during the same time period.
Blinding	Probably low risk	Study reports that samples were coded, which suggests blinding measures were implemented.
Exposure Assessment	Low risk	The method of alpha particle concentration assessment (CR-39 fission track detector) was robust.
Outcome Assessment	Probably low risk	The authors do not sufficiently describe how leukemia patients were diagnosed, but the use of the term 'patients' suggests that they were diagnosed by medical professionals.
Confounding	High risk	The study collected data on age and sex of participants, but did not control for those or any other confounders.
Incomplete Outcome Data	Low risk	Study is free of missing outcome data.
Selective Reporting	Low risk	All pre-specified outcomes were reported.
Other Bias	Low risk	Study appears free of other sources of bias.
Conflict of Interest	Probably low risk	Associated funds and persons appear to be from academia only and free of financial interests in study results. However, no claim denying conflicts of interest was made.

Note: For more information, please see Supplemental Material, "The Navigation Guide instructions for making risk of bias determinations"

Data abstraction items

Study Characteristic Items	Risk of Bias Items
Target population	Recruitment strategy for participants
Total number of study participants	Recruitment strategy for comparator/controls
Characteristics of participants (age, gender, etc.)	Response Rate
Type of exposure assessed	Blinding measures in place
Characteristics of comparator/controls	Confounding variables measured/assessed
Type of health outcome assessed	Exposure assessment methods
Method of health outcome assessment	Time of exposure measurement
Time of health outcome measurement	Location of exposure measurement
Year of publication	Strategy for addressing missing outcome data
Study setting	Evidence of selective outcome reporting
Study design	Other sources of methodological bias
Publication status	Financial conflict of interest
Association measured	
Major finding/conclusions of study	
Notes	

Screening form

Does the publication report or measure uranium, its corrosion products, or ionizing radiation in Iraq or surrounding areas?

Uranium corrosion products may include uranium oxides and uranyl ions, while exposure to ionizing radiation may include alpha, beta, or gamma rays. Studies that only measure or report exposure to UV radiation (i.e. solar radiation) are excluded. Levels or concentrations of ionizing radiation or uranium should be measured/reported in Iraq or surrounding areas such as Kuwait.

- NO → Exclude, and note reason in Excel
 Yes → Go to the next question

Does the publication report or measure health outcomes or disease states in humans?

Do not restrict to only birth outcomes at this stage. Outcomes ought to be clinical or “patient important”, i.e. exclude if they only measure uranium concentrations in human tissues.

- NO → Exclude, and note reason in Excel
 Yes → Go to the next question

Is the study population Iraqi?

Does the human population for which health outcomes are reported include or consist entirely of Iraqi nationals? Exclude studies that only report health outcomes in populations of military veterans (of the US or other nationalities) who fought during the 1990 or 2003 Iraq Wars.

- NO → Exclude, and note reason in Excel
 Yes → Go to the next question

Does the publication include or report primary research?

Exclude review articles, including systematic reviews, which do not contain or report primary research.

- NO → Exclude, and note reason in Excel
 Yes → Go to the next question

Does the study include a nonexposed (or lesser exposed) comparator or control group?

Exclude single-arm, non-comparator studies.

- NO → Exclude, and note reason in Excel
 Yes → INCLUDE

The Navigation Guide instructions for making risk of bias determinations

1. Was the strategy for recruiting participants consistent across study groups?

Criteria for a judgment of 'YES' (i.e. low risk of bias):

Protocols for recruitment and inclusion/exclusion criteria were applied similarly across study groups, and any one of the following:

- Study participants were recruited from the same population at the same time frame; or
- Study participants were not all recruited from the same population, but proportions of participants from each population in each study group are uniform

Criteria for the judgment of 'PROBABLY YES' (i.e. probably low risk of bias):

There is insufficient information about participant selection to permit a judgment of 'YES', but there is indirect evidence that suggests that participant recruitment and inclusion/exclusion criteria was consistent, as described by the criteria for a judgment of 'YES'.

Criteria for the judgment of 'NO' (i.e. high risk of bias)

Any one of the following:

- Protocols for recruitment or inclusion/exclusion criteria were applied differently across study groups; or
- Study participants were recruited at different time frames; or
- Study participants were recruited from different populations and proportions of participants from each population in each study group are not uniform

Criteria for the judgment of 'PROBABLY NO' (i.e. probably high risk of bias):

There is insufficient information about participant selection to permit a judgment of 'NO', but there is indirect evidence that suggests that participant recruitment or inclusion/exclusion criteria was inconsistent, as described by the criteria for a judgment of 'NO'.

Criteria for the judgment of 'NOT APPLICABLE' (risk of bias domain is not applicable to study):

There is evidence that participant selection is not an element of study design capable of introducing risk of bias in the study.

2. Was knowledge of the exposure groups adequately prevented during the study?

Criteria for a judgment of 'YES' (i.e. low risk of bias):

Any one of the following:

- No blinding, but the review authors judge that the outcome and the outcome measurement are not likely to be influenced by lack of blinding; or
- Blinding of key study personnel ensured, and unlikely that the blinding could have been broken; or
- Some key study personnel were not blinded, but outcome assessment was blinded and the non-blinding of others unlikely to introduce bias.

Criteria for the judgment of ‘PROBABLY YES’ (i.e. probably low risk of bias):

There is insufficient information about blinding to permit a judgment of ‘YES’, but there is indirect evidence that suggests the study was adequately blinded, as described by the criteria for a judgment of ‘YES’.

Criteria for the judgment of ‘NO’ (i.e. high risk of bias):

Any one of the following:

- No blinding or incomplete blinding, and the outcome or outcome measurement is likely to be influenced by lack of blinding; or
- Blinding of key study personnel attempted, but likely that the blinding could have been broken; or
- Some key study personnel were not blinded, and the non-blinding of others likely to introduce bias.

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

There is insufficient information about blinding to permit a judgment of ‘NO’, but there is indirect evidence that suggests the study was not adequately blinded, as described by the criteria for a judgment of ‘NO’.

Criteria for the judgment of ‘NOT APPLICABLE’ (risk of bias domain is not applicable to study):

There is evidence that blinding is not an element of study design capable of introducing risk of bias in the study.

3. Were exposure assessment methods robust?

Criteria for a judgment of ‘YES’ (i.e. low risk of bias):

The reviewers judge that there is low risk of exposure misclassification and any one of the following:

- There is high confidence in the accuracy of the exposure assessment methods; or
- Less-established or less direct exposure measurements are validated against well-established or direct methods AND if applicable, appropriate QA/QC for methods are described and are satisfactory, with at least three of the following items reported, or at least two of the following items reported plus evidence of

satisfactory performance in a high quality inter-laboratory comparison: Limit of detection or quantification; standards recovery; measure of repeatability; investigation and prevention of blanks contamination.

Criteria for the judgment of ‘PROBABLY YES’ (i.e. probably low risk of bias):

There is insufficient information about the exposure assessment methods to permit a judgment of ‘YES’, but there is indirect evidence that suggests that methods were robust, as described by the criteria for a judgment of ‘YES’. Studies only reporting that the QA/QC items above were satisfactory but not reporting all of the actual numbers may receive a judgment of “probably yes.”

Criteria for the judgment of ‘NO’ (i.e. high risk of bias):

The reviewers judge that there is high risk of exposure misclassification and any one of the following:

- There is low confidence in the accuracy of the exposure assessment methods; or
- Less-established or less direct exposure measurements are not validated and are suspected to introduce bias that impacts the outcome assessment (example: participants are asked to report exposure status retrospectively, subject to recall bias)
- Uncertain how exposure information was obtained

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

There is insufficient information about the exposure assessment methods to permit a judgment of ‘NO’, but there is indirect evidence that suggests that methods were not robust, as described by the criteria for a judgment of ‘NO’.

Criteria for the judgment of ‘NOT APPLICABLE’ (risk of bias domain is not applicable to study):

There is evidence that exposure assessment is not an element of study design capable of introducing risk of bias in the study.

4. Were outcome assessment methods robust?

Criteria for a judgment of ‘YES’ (i.e. low risk of bias):

The reviewers judge that there is low risk of outcome misclassification, i.e.:

- Outcomes were assessed and defined consistently across all study participants, using valid and reliable measures; or
- Less-established or less direct outcome measurements are validated against well-established or direct methods; or
- Appropriate sensitivity analyses were conducted that suggest the influence of outcome misclassification would be minimal

- AND, if applicable, appropriate QA/QC for methods is described and is satisfactory.

Criteria for the judgment of ‘PROBABLY YES’ (i.e. probably low risk of bias):

There is insufficient information about the outcome assessment methods to permit a judgment of low risk of bias, but there is indirect evidence which suggests that methods were robust, as described by the criteria for a judgment of low risk of bias. Appropriate QA/QC for methods are not described but the review authors judge that the outcome and the outcome assessment are objective and uniform across study groups.

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

There is insufficient information about the outcome assessment methods to permit a judgment of high risk of bias, but there is indirect evidence which suggests that methods were not robust, as described by the criteria for a judgment of high risk of bias.

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

The reviewers judge that there is high risk of outcome misclassification and any one of the following:

- There is low confidence in the accuracy of the outcome assessment methods; or
- Less-established or less direct outcome measurements are not validated and are suspected to introduce bias that impacts the outcome assessment
- Uncertain how outcome information was obtained

Criteria for the judgment of ‘NOT APPLICABLE’ (risk of bias domain is not applicable to study):

There is evidence that outcome assessment methods are not capable of introducing risk of bias in the study.

5. Was confounding adequately addressed?

Criteria for a judgment of ‘YES’ (i.e. low risk of bias):

The study accounted for (i.e., matched, stratified, multivariate analysis or otherwise statistically controlled for) important potential confounders, or reported that potential confounders were evaluated and omitted because inclusion did not substantially affect the results. The determination of specific confounders may be informed by the data, including the studies included in the review.

Criteria for the judgment of ‘PROBABLY YES’ (i.e. probably low risk of bias):

The study accounted for most but not all of the important potential confounders AND this lack of accounting is not expected to introduce substantial bias.

Criteria for the judgment of 'NO' (i.e. high risk of bias):

The study did not account for or evaluate important potential confounders.

Criteria for the judgment of 'PROBABLY NO' (i.e. probably high risk of bias):

The study accounted for some but not all of the important potential confounders AND this lack of accounting may have introduced substantial bias.

6. Were incomplete outcome data adequately addressed?

Criteria for a judgment of 'YES' (i.e. low risk of bias):

Participants were followed long enough to obtain outcome measurements and any one of the following:

- No missing outcome data; or
- Reasons for missing outcome data unlikely to be related to true outcome (for survival data, censoring unlikely to be introducing bias); or
- Missing outcome data balanced in numbers across exposure groups, with similar reasons for missing data across groups; or
- For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk not enough to have a biologically relevant impact on the intervention effect estimate; or
- For continuous outcome data, plausible effect size (difference in means or standardized difference in means) among missing outcomes not enough to have a biologically relevant impact on observed effect size; or
- Missing data have been imputed using appropriate methods

Criteria for the judgment of 'PROBABLY YES' (i.e. probably low risk of bias):

There is insufficient information about incomplete outcome data to permit a judgment of 'YES', but there is indirect evidence that suggests incomplete outcome data was adequately addressed, as described by the criteria for a judgment of 'YES'.

Criteria for the judgment of 'NO' (i.e. high risk of bias):

Participants were not followed long enough to obtain outcome measurements OR any one of the following:

- Reason for missing outcome data likely to be related to true outcome, with either imbalance in numbers or reasons for missing data across exposure groups; or

- For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk enough to induce biologically relevant bias in intervention effect estimate; or
- For continuous outcome data, plausible effect size (difference in means or standardized difference in means) among missing outcomes enough to induce biologically relevant bias in observed effect size; or
- Potentially inappropriate application of imputation.

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

There is insufficient information about incomplete outcome data to permit a judgment of ‘NO’, but there is indirect evidence that suggests incomplete outcome data was not adequately addressed, as described by the criteria for a judgment of ‘NO’.

Criteria for the judgment of ‘NOT APPLICABLE’ (risk of bias domain is not applicable to study):

There is evidence that incomplete outcome data is not capable of introducing risk of bias in the study.

7. Are reports of the study free of suggestion of selective outcome reporting?

Criteria for a judgment of ‘YES’ (i.e. low risk of bias):

All of the study’s pre-specified (primary and secondary) outcomes outlined in the protocol, methods, abstract, and/or introduction that are of interest in the review have been reported in the pre-specified way.

Criteria for the judgment of ‘PROBABLY YES’ (i.e. probably low risk of bias):

There is insufficient information about selective outcome reporting to permit a judgment of ‘YES’, but there is indirect evidence that suggests the study was free of selective reporting, as described by the criteria for a judgment of ‘YES’.

Criteria for the judgment of ‘NO’ (i.e. high risk of bias):

Any one of the following:

- Not all of the study’s pre-specified primary outcomes (as outlined in the protocol, methods, abstract, and/or introduction) have been reported; or
- One or more primary outcomes is reported using measurements, analysis methods or subsets of the data (e.g. subscales) that were not pre-specified; or
- One or more reported primary outcomes were not pre-specified (unless clear justification for their reporting is provided, such as an unexpected effect); or
- One or more outcomes of interest are reported incompletely

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

There is insufficient information about selective outcome reporting to permit a judgment of 'NO', but there is indirect evidence that suggests the study was not free of selective reporting, as described by the criteria for a judgment of 'NO'.

Criteria for the judgment of 'NOT APPLICABLE' (risk of bias domain is not applicable to study):

There is evidence that selective outcome reporting is not capable of introducing risk of bias in the study.

8. Was the study apparently free of other problems that could put it at a risk of bias?

Criteria for a judgment of 'YES' (i.e. low risk of bias):

The study appears to be free of other sources of bias.

Criteria for the judgment of 'PROBABLY YES' (i.e. probably low risk of bias):

There is insufficient information to permit a judgment of 'YES', but there is indirect evidence that suggests the study was free of other threats to validity.

Criteria for the judgment of 'NO' (i.e. high risk of bias):

There is at least one important risk of bias. For example, the study:

- Had a potential source of bias related to the specific study design used; or
- Stopped early due to some data-dependent process (including a formal-stopping rule); or
- Had extreme imbalance of characteristics among exposure groups; or
- Had differential surveillance for outcome between exposure groups or between exposed/unexposed groups
- The conduct of the study is affected by interim results (e.g. recruiting additional participants from a subgroup showing greater or lesser effect); or
- An insensitive instrument is used to measure outcomes (which can lead to under-estimation of both beneficial and harmful effects); or
- Selective reporting of subgroups; or • Has been claimed to have been fraudulent; or
- Had some other problem

Criteria for the judgment of 'PROBABLY NO' (i.e. probably high risk of bias):

There is insufficient information to permit a judgment of 'NO', but there is indirect evidence that suggests the study was not free of other threats to validity, as described by the criteria for a judgment of 'NO'.

Criteria for the judgment of 'NOT APPLICABLE' (risk of bias domain is not applicable to study):

There is evidence that other potential threats to validity are not capable of introducing risk of bias in the study.

9. Was the study free of support from a company, study author, or other entity having a financial interest in any of the exposures studied?

Criteria for a judgment of 'YES' (i.e. low risk of bias):

The study did not receive support from a company, study author, or other entity having a financial interest in the outcome of the study. Examples include the following:

- Funding source is limited to government, non-profit organizations, or academic grants funded by government, foundations and/or non-profit organizations;
- Chemicals or other treatment used in study were purchased from a supplier;
- Company affiliated staff are not mentioned in the acknowledgements section;
- Authors were not employees of a company with a financial interest in the outcome of the study;
- Company with a financial interest in the outcome of the study was not involved in the design, conduct, analysis, or reporting of the study and authors had complete access to the data;
- Study authors make a claim denying conflicts of interest;
- Study authors are unaffiliated with companies with financial interest, and there is no reason to believe a conflict of interest exists;
- All study authors are affiliated with a government agency (are prohibited from involvement in projects for which there is a conflict of interest or an appearance of conflict of interest).

Criteria for the judgment of 'PROBABLY YES' (i.e. probably low risk of bias):

There is insufficient information to permit a judgment of 'YES', but there is indirect evidence that suggests the study was free of support from a company, study author, or other entity having a financial interest in the outcome of the study, as described by the criteria for a judgment of 'YES'.

Criteria for the judgment of 'NO' (i.e. high risk of bias):

The study received support from a company, study author, or other entity having a financial interest in the outcome of the study. Examples of support include:

- Research funds;
- Chemicals provided at no cost;
- Writing services;
- Author/staff from study was employee or otherwise affiliated with company with financial interest;

- Company limited author access to the data;
- Company was involved in the design, conduct, analysis, or reporting of the study;
- Study authors claim a conflict of interest

Criteria for the judgment of ‘PROBABLY NO’ (i.e. probably high risk of bias):

There is insufficient information to permit a judgment of ‘NO’, but there is indirect evidence that suggests the study was not free of support from a company, study author, or other entity having a financial interest in the outcome of the study, as described by the criteria for a judgment of ‘NO’.

Criteria for the judgment of ‘NOT APPLICABLE’ (risk of bias domain is not applicable to study):

There is evidence that conflicts of interest are not capable of introducing risk of bias in the study.

Full text exclusion justifications

Include Report met all inclusion criteria, as specified in the PECOS statement	[1-33]
Exclude – Exposure Study did not measure or report exposure to weaponized uranium in Iraq	[34-74]
Exclude – Outcome Study did not measure or report patient-important health outcomes	[75-121]
Exclude – Population Study population was not Iraqi	[122-147]
Exclude – Study Design: Primary research Study did not contain original data or analysis	[148-203]
Exclude – Study Design: Comparator Non-comparator study	[204-225]

Include

1. Abdul-Wahid R. 2009. Depleted uranium effects on immunophenotyping of human lymphocytes in southern Iraq. *Iraqi Journal of Community Medicine* 22(4):249-253.
2. Al-Hamadany WS, Saleh DS, Shanshal MA. 2012. Radiation pollution in cancer and other diseases using some immunological and clinical parameters. *The Iraqi Journal of Veterinary Medicine* 36(2):33-40.
3. Al-Hamzawi AA, Jaafar MS, Tawfiq NF. 2014. Uranium concentration in blood samples of southern Iraqi leukemia patients using CR-39 track detector. *Journal of Radioanalytical and Nuclear Chemistry* 299:1267-1272. PMID: 26224958, <https://doi.org/10.1007/s10967-013-2808-0>.
4. Al-Hamzawi AA, Jaafar MS, Tawfiq NF. 2015. Concentration of uranium in human cancerous tissues of southern Iraqi patients using fission track analysis. *Journal of Radioanalytical and Nuclear Chemistry* 303:1703-1709. <https://doi.org/10.1007/s10967-014-3682-0>.

5. Al-Hashimi MMY, Wang X. 2013. Comparing the cancer in Ninawa during three periods (1980-1990, 1991-2000, 2001-2010) using poisson regression. *Journal of Research in Medical Sciences* 18(12):1026-1039. PMID: 24523792.
6. Al-Jobori S. 2013. Track detection technique using cr-39 for determining depleted uranium in biological specimens. *Journal of Medenat Alelem College* 5(2):5-14.
7. Al-Rudainy LA, Ajeel NA, Al-Saad HT. 2009a. Depleted uranium and incidence of cancer in Basrah: A preliminary ecological study. *The Medical Journal of Basrah University* 27(1):1-6.
8. Al-Rudainy LA, Salih HM, Aldorky MK. 2009b. Incidence and pattern of childhood leukaemia in Basrah, Iraq during 2003– 2007. *Iran Journal of Blood and Cancer* 2(1):11-17.
9. Al-Sabbak M, Sadik Ali S, Savabi O, Savabi G, Dastgiri S, Savabieasfahani M. 2012. Metal contamination and the epidemic of congenital birth defects in Iraqi cities. *Bulletin of Environmental Contamination and Toxicology* 89(5):937-944. PMID: 22983726, <https://doi.org/10.1007/s00128-012-0817-2>.
10. Al-Sadoon I, Hassan GG, Yacoub AA. 1999. Depleted uranium and health of people in Basrah: Epidemiological evidence. 2. The incidence and pattern of congenital anomalies among births in Basrah during the period 1990–1998. *The Medical Journal of Basrah University* 17:27-33.
11. Al-Sahlane MHR., Ramli RM, Ali MAH, Tawfiq N., Rahman AA, Mustafa IS, Azman NZN, Razak N, Yahaya NZ, Al-Marri HM, Ayob NS, Zakaria N. 2016. Analysis of Uranium concentration on maternal and umbilical cord blood samples after delivery in Iraq. 6th IEEE International Conference on Control System, Computing and Engineering (ICCSCE) (pp. 360-365). IEEE. <https://doi.org/10.1109/ICCSCE.2016.7893599>.
12. Al-Sahlane MHR., Ramli RM, Ali MAH, Tawfiq N., Rahman AA, Mustafa IS, Azman NZN, Razak N, Yahaya NZ, Al-Marri HM, Ayob NS, Zakaria N. 2017. Trace of heavy metals in maternal and umbilical cord blood samples in association with birth outcomes in Baghdad, Iraq. *Proceedings of the Regional Conference on Nuclear Physics, RCNP. EPJ Web of Conferences* 156:00003. <https://doi.org/10.1051/epjconf/201715600003>.
13. Alaani S, Tafash M, Busby C, Hamdan M, Blaurock-Busch E. 2011. Uranium and other contaminants in hair from the parents of children with congenital anomalies in Fallujah, Iraq. *Conflict and Health* 5:15. PMID: 21888647, <https://doi.org/10.1186/1752-1505-5-15>.
14. Alaani S, Al-Fallouji M, Busby C, Hamdan M. 2012. Pilot study of congenital anomaly rates at birth in Fallujah, Iraq, 2010. *Journal of the Islamic Medical Association of North America* 44(1):1-7. PMID: 23864991, <https://www.doi.org/10.5915/44-1-10463>.
15. Alborz A. 2013. Environmental characteristics and prevalence of birth defects among children in post-war iraq: Implications for policies on rebuilding the iraqi education system. *Medicine, Conflict and Survival* 29(1):26-44. PMID: 23729096, <https://www.doi.org/10.1080/13623699.2013.765197>.

16. Al-Rudainy LA, Hassan JG, Salih HM, Abbas MK, Majeed AA. 2011. Time trends and geographical distribution of childhood leukaemia in Basrah, Iraq, from 2004 to 2009. *Sultan Qaboos University Medical Journal* 11(2):215-220. PMID: 21969893.
17. Busby C, Hamdan M, Ariabi E. 2010. Cancer, infant mortality and birth sex-ratio in Fallujah, Iraq 2005-2009. *International Journal of Environmental Research and Public Health* 7(7):2828-2837. PMID: 20717542, <https://www.doi.org/10.3390/ijerph7072828>.
18. Hagopian A, Lafta R, Hassan J, Davis S, Mirick D, Takaro T. 2010. Trends in childhood leukemia in Basrah, Iraq, 1993–2007. *American Journal of Public Health* 100(6):1081-1087. PMID: 20167894, <https://www.doi.org/10.2105/AJPH.2009.164236>.
19. Hassan JKA, N. A. H., Hamadi SS. 2005. Incidence and time trend of cancer in Basrah. *The Medical Journal of Basrah University* 23(2):13-20.
20. Humadi A, Khalaf S. 2011. Hematological and cytogenetic study for person shot by coalition forces in al-anbar governorate. *Tikrit Journal of Pure Science* 16:92-99.
21. Mryoush AQ, Salim HM. 2015. Determination of uranium concentration in soil of baghdad governorate and its effect on mitotic index assay. *Iraqi Journal of Science* 56:140-146.
22. Neamah KA, Tawfiq AA. 2015. Analysis of pollution in al-fallujah district by using estimation of autoregressive coefficients. *Al-Ma'mon College Journal*:237-247.
23. Mohammad FI. 2016. Tissue microarray construction and immunohistochemical evaluation of bcl-2 gene expression in iraqi and italian breast cancer samples *Jornal of Biotechnology Research Center* 10:32-38.
24. Salman SM. 2008. Study the effect of depleted uranium used by coalition forces in increasing of cancer disease in diyala governorate. *Diyala Journal of Human Research*:144-149.
25. Savabieasfahani MA, S. S.; Bacho, R.; Savabi, O.; Alsabbak, M. 2016. Prenatal metal exposure in the middle east: Imprint of war in deciduous teeth of children. *Environmental Monitoring and Assessment* 188.
26. Shafik SS. 2014. Study and measurements of the uranium and amorphous crystals concentrations in urine samples of breast cancer female patients. *Iraqi Journal of Physics* 12:113-122.
27. Abed, M.M., K.H. Mahdi, and W.S. Al-Hamadany. Estimation of uranium concentration in blood samples of kidneys failure patients in Al-Muthanna governorate. 2019.
28. Hamadany, W., H.M. Khalid, and M.A. Muayad, Estimation of Uranium Concentration in Urine Samples of Kidneys Failure Patients in Al- Muthanna Governorate. *Ibn Al-Haitham Journal For Pure And Applied Science* *مجلة ابن الهيثم للعلوم والتطبيقية الصرفة للعلوم*, 2020. 33(1): p. 11-16.
29. Hassan, A.B., et al., Determination of alpha particles levels in blood samples of cancer patients at Karbala Governorate, Iraq. *Iranian Journal of Medical Physics*, 2019. 16(1): p. 41-47.

30. Jumaah, A.S., et al., Increased PTEN gene expression in patients with endometrial carcinoma from areas of high risk depleted uranium exposure. *BMC Res Notes*, 2019. 12(1): p. 708.
31. Qaddoori, S.M. and S.S. Shafik, CR-39 as a tool for uranium concentration calculation in bio assay sample: Bladder cancer as case study. *Research Journal of Pharmaceutical, Biological and Chemical Sciences*, 2018. 9(6): p. 228-239.
32. Savabieasfahani, M., F. Basher Ahamadani, and A. Mahdavi Damghani, Living near an active U.S. military base in Iraq is associated with significantly higher hair thorium and increased likelihood of congenital anomalies in infants and children. *Environ Pollut*, 2020. 256: p. 113070.
33. Showard, A.F. and M.S. Aswood. Measuring of Alpha particles in Blood samples of Leukemia patients in Babylon governorate, Iraq. 2019.

Exclude – Exposure

34. Al-Zacko, S.M.Z., H. G.; Mohammad, A. S., *Pediatric burns in Mosul: an epidemiological study*. *Annals of Burns & Fire Disasters*. **27**(2): p. 70-5.
35. Gousheh, J.R., M., [*War injuries of the femoral nerve. Apropos of a series of 27 cases*]. *Annales de Chirurgie Plastique et Esthetique*. **36**(6): p. 527-31.
36. Balasem, A.N.A., A. Sk; Hs.; Hussain, K. O., *Chromosomal Aberration Analysis in Peripheral Lymphocytes of Radiation Workers*. *Mutation Research*, 1992. **271**(3): p. 209-211.
37. Wareham, S., *IPPNW, 1991: Disarmament, development and the environment*. *Medical Journal of Australia*, 1992. **156**(7): p. 443-444.
38. Luca, D., *Intervention humanitaire: Questions et reflexions*. *International Journal of Refugee Law*, 1993. **5**(3): p. 424-441.
39. Weimaster, J.F.B., W. T.; Bossle, P. C.; Ellzy, M. W.; Janes, L. G.; Johnson, D. W.; Lochner, J. M.; Pleva, S. G.; Reeder, J. H.; Rohrbaugh, D. K.; Rosso, T. E.; Szafraniec, L. J.; Szafraniec, L. L.; Albro, T. G.; Creasy, W. R.; Stuff, J. R.; Smith, P. B.; Stewart, I. R., *Chemical analysis of environmental samples collected in iraq analysis for the presence of chemical warfare agents*. *Journal of Chemical Technology and Biotechnology*, 1995. **64**(2): p. 115-128.
40. Charp, P.A., *Al Eskan disease: Persian Gulf syndrome*. *Military medicine*, 1997. **162**(3).
41. Camus, Y.L.V., C.; Louvart, L., *GOGP (Gulf of Oman, Persian Gulf): SHOM's oceanography campaign*. *Annales Hydrographiques*, 2002. **2**(771): p. 31-36.
42. Nassir, T.H.A.-A., S. M., *A retrospective study of head and neck malignancies, in International Congress Series*. 2003. p. 1067-1074.
43. Schmid, C.R., J. S., *Neurodevelopmental toxicology*. *Neurologic Clinics*, 2005. **23**(2): p. 321-336.
44. Peplow, M., *Counting the dead*. *Nature*, 2006. **440**(7087): p. 982-983.

45. Dr. Sattar Jabbar جبار سنار دكتور ا.ا., *Variables in the Iraqi market and its impact on consumer protection in the future*. Iraqi Journal For Economic Sciences العراقية المجلة للاقتصادية للعلوم, 2007(13): p. 69-85.
46. Abouzaid, H., *Health and the environment with focus on the Eastern Mediterranean Region*. EMHJ - Eastern Mediterranean Health Journal, 2008. **14**(Special Issue): p. S132-S142.
47. Lavin, R.P., *Histologic Changes as Indicators of Carcinogenicity of Tungsten Alloy in Rodents*. 2008.
48. Nasser abed saggad عبد نصر ا.ا., *Journals geographic*, 2008. **1**(10): p. 34-45.
49. Ismail, A.H.J., M. S., *Relationship between radon concentration, ventilation rate and male infertility: A case study in Iraqi Kurdistan*. International Journal of Low Radiation, 2010. **7**(3): p. 175-187.
50. Sallama, S.H.N., F. Tawfiq, *DETERMINATION OF URANIUM CONCENTRATION IN TEETH MALESAMPLES USING FISSION TRACKS IN CR-39 FROM DIFFERENTCOUNTRIES*. Journal of Al-Nahrain University - Science مجلة كلية العلوم - جامعة - النهرين جامعة, 2010. **13**(1): p. 91-95.
51. Guillemin, D., *The navy and the gulf war*. Guerres Mondiales et Conflicts Contemporains, 2011. **244**(4): p. 31-51.
52. Mousavi, S.M.B., A.; Sundquist, J.; Hemminki, K., *Risks of papillary and follicular thyroid cancer among immigrants to Sweden*. International Journal of Cancer, 2011. **129**(9): p. 2248-2255.
53. Ganjo, *20th International Conference on Modelling, Monitoring and Management of Air Pollution, AIR12*. 20th International Conference on Modelling, Monitoring and Management of Air Pollution, AIR 2012, 2012. **157**.
54. Mateen, F.J.C., M.; Al-Saedy, H.; Nyce, S.; Ghosn, J.; Mutuerandu, T.; Black, R. E., *Medical conditions among Iraqi refugees in Jordan: Data from the United Nations refugee assistance information system*. Bulletin of the World Health Organization, 2012. **90**(6): p. 444-451.
55. Protopsaltis, *20th International Conference on Modelling, Monitoring and Management of Air Pollution, AIR12*. 20th International Conference on Modelling, Monitoring and Management of Air Pollution, AIR 2012, 2012. **157**.
56. Strain, J.W., 2nd, *CBRNE TC3: A Hybrid Approach to Casualty Care in the CBRNE Environment*. Journal of special operations medicine : a peer reviewed journal for SOF medical professionals, 2013. **13**(2): p. 44-53.
57. حمادي, A.A.H.ع.ا.ع.ا.ا., *A study of some Biological and Biochemical Indicators to Al-Falluja Citizens after the Second Invasion of the City by Multi-National Forces*. Al-Anbar Journal of Veterinary Sciences البيطرية للعلوم الانبار مجلة, 2013. **6**(1): p. 248-261.
58. Al-Shammari, A.M.A.-J., A. A.; Al-Mukhtar, A. A.; Ali, A. M.; Al-Hili, Z. A.; Yaseen, N. Y., *Establishment and characterization of a chemoresistant glioblastoma cell line from an Iraqi patient*. Cancer Research, 2014. **74**(19).
59. Faisal, N.N.ن.ن.ف.ن.ن., *The use of cluster analysis to classify cancer by species And situations in Iraq for the period (1998 - 2006) (1998 – 2006)*. Journal of kirkuk University For Administrative and Economic Sciences, 2014. **4**(1): p. 269-291.

60. Biko, D.M.M., B. F.; Jesinger, R. A.; Sherman, P. M.; Borg, B. D.; Lichtenberger, J. P., *Imaging of pediatric pathology during the Iraq and Afghanistan conflicts*. Pediatric Radiology, 2015. **45**(3): p. 439-448.
61. Jaffar, S., *La politique culturelle du Gouvernement Régional du Kurdistan - Irak*. Maghreb - Machrek, 2015. **222**(4): p. 27-43.
62. Virof, E.S., E.; Laurent, F.; Ferry, T.; Lyon, Bone; Joint Infection Study, Group, *Reactivation of Clostridium tertium bone infection 30 years after the Iran-Iraq war*. BMJ Case Reports, 2015. **18**: p. 18.
63. Anfal Saied Dawood سعيد انفال د.س.أ.م.س.و.ع.س.منخي, *Environmental radioactive contamination and its effects on the spread of cancer in the province of Baghdad (Amal Hospital model)* (Al-Adab Journal, 2016(119): p. 375-392.
64. Patel, M.N., *Neuroprotective effects of AEOL 10150 against organophosphate toxicity*, in *RePORTER Database National Institutes of Health*. 2016.
65. Herzmann, C.G., M.; Malekzada, F.; Lonroth, K.; Kranzer, K., *Radiological screening of refugees in Germany*. European Respiratory Journal, 2017. **49**(5).
66. Jabir, F.A.H., W. H., *No Evaluation of Serum P53 Levels in Iraqi Female Breast Cancer Patients*. Asian Pac J Cancer Prev, 2017. **18**(9): p. 2551-2553.
67. Abbas Kadhum, R. and W. Abbas Aldrghi. *Detection of XRCC1 Expression and (8-OHdG) Levels as a marker of Oxidative DNA Damage in Individuals Exposed to Low Dose of Gamma Rays*. 2019.
68. Aldin, S.Z.J., et al., *Work-related and prostate cancer of man and woman in Mosul/Iraq*. International Journal of Psychosocial Rehabilitation, 2020. **24**(4): p. 1729-1736.
69. Al-Drighi, W.A. and M.T. Hassan, *Molecular study of ahr gene polymorphisms among iraqi women staff working in radiotherapy units*. International Journal of Pharmaceutical Quality Assurance, 2019. **10**(2): p. 285-289.
70. Al-Jameel, L.S., R.S. Ahmed, and H.A. Shamran, *Ionizing radiation effect and DNA damage in the workers of Al-Tuwaitha nuclear site*. Iraqi Journal of Science, 2019. **60**(12): p. 2636-2641.
71. Al-Mukhtar, S., et al., *Relation between osteoarthritis and HLA-A in Iraqi patients*. Prensa Medica Argentina, 2018. **104**(5): p. 261-264.
72. Aziz, H.A., *Effect of radiation on employers in radiology (X- rays, CT Scan and Magnetic Resonance Imaging)In Al-muthanna Province-Iraq*. Journal of Global Pharma Technology, 2018. **10**(3): p. 981-984.
73. Ouda, Z.A., A.N. Kaloub, and A.K. Ali, *Assessment of dna damage in lymphocytes of radiationworkers at al-tuwaitha site using nuclear division index and hypoxanthine guanine phosphoribosyl transferase (HPRT)*. Research Journal of Pharmacy and Technology, 2018. **11**(7): p. 2855-2858.
74. Rasheed, Z.A., A.K. Ali, and A.A. Majed, *Evaluation of micronuclei in radiationworkers chronically exposed to low dose ionizing radiation*. Annals of Tropical Medicine and Public Health, 2019. **22**(Special Issue 5).

Exclude – Outcome

75. *Facing the nuclear danger*. Nature, 2003. **426**(6964): p. 213.
76. *Worldwide Emerging Environmental Issues Affecting the U.S. Military. October 2008 Report*. Govt Reports Announcements & Index, 2008(26): p. 21.
77. *Support for environmental management of the Iraqi marshlands 2004-2009*. 2009, United Nations Environment Programme.
78. Abojassim, A.A.M., H. A. A.; Husain, A. A.; Wood, M., *Estimation of the excess lifetime cancer risk from radon exposure in some buildings of Kufa technical institute, Iraq*. Nuclear Physics and Atomic Energy, 2017. **18**(3): p. 276-286.
79. AGENCY, I.A.E., *Radiological Conditions in Selected Areas of Southern Iraq with Residues of Depleted Uranium*. 2010, Vienna: INTERNATIONAL ATOMIC ENERGY AGENCY.
80. Al Kinani, A.A.S., AS; Al-Anni, S, *Investigation of depleted uranium contamination in south west of Iraq*. 2005.
81. Al-Ghais, F.R., *Communication dated 27 January 1994 received from the resident representative of Kuwait to the International Atomic Energy Agency referring to document INFCIRC/425*. Govt Reports Announcements & Index, 1994(06): p. 18.
82. Al-Hamzawi, A.A.J., M. S.; Tawfiq, N. F., *The measurements of uranium concentration in human blood in selected regions in Iraq using CR-39 track detector*, in *Joint International Conference on Nanoscience, Engineering and Management, BOND21*. 2014, Trans Tech Publications: Penang. p. 679-683.
83. Al-Jobouri, A.F.S., *Determination of Uranium Concentration in Human Urine for Selected Regions in Iraq Using Laser-Induced Kinetic Phosphorimetry and CR-39 Nuclear Track Detector*. 2012, MSc dissertation, Dept. Phys. College of Science, Al-Nahrain Univ., Iraq.
84. Al-Lami, A.G., H.; Abdulhamid, H.; Mukhtar, E.; Hilo, T., *The National Environmental Strategy and Action Plan for Iraq (2013 – 2017)*. 2013, Iraqi Ministry of Environment.
85. Almayahi, B., *Determination of radionuclide concentration in human teeth in Najaf governorate, Iraq*. Iranian Journal of Medical Physics, 2017. **14**(4): p. 173-182.
86. Almayahi, B.H., J. I.; Saheb, L., *The impact of low-level exposure to radiation in natural ecosystems of Najaf and Dhi Qar Cities, Iraq*. Iranian Journal of Medical Physics, 2018. **15**(1): p. 1-5.
87. Alya'a Abdurazak عبد علياء, ا.ن., F. Tawfiq توفيق فاضل ندى; Nidhala, H. Kadhim نضالة كاظم حسن, *Determination of Uranium Concentration in child teeth by track detector CR-39 for middle and south of Iraq*. Baghdad Science Journal, 2010. **7**(1): p. 1-5.
88. Alya'a Abdurazak عبد علياء, ا.ن., F. Tawfiq توفيق فاضل ندى; Nidhala, H. Kadhim نضالة كاظم حسن, *Determination of Uranium Concentration in child teeth by track detector CR-39 in same middle and south regions of Iraq CR-39*. Baghdad Science Journal مجلة بغداد للعلوم, 2011. **8**(4): p. 909-913.
89. Brand, R.A.S., E., *Depleted uranium: A new environmental radiotoxicological pollutant*. Landbauforschung Volkenrode, 2005. **55**(4): p. 211-218.
90. Eick, R., N. Khalil, and P. Blair, *CHILDHOOD EXPOSURE TO DEPLETED URANIUM: CONSEQUENCES OF THE 2003 ARMED CONFLICT IN IRAQ*.
91. Gerdes, A.W., S; Brey, G; Zimmermann, I; Durakovic, A, *Precise Monitoring of Depleted Uranium in human and environment of South Iraq using Multi-collector ICP-MS*.

92. Israa, K.A. الك.ا.ا., A. Razzaq عبد علياء الرزاق, *Measuring the concentration of uranium for adults teeth in adjacent areas of Tigris river in Baghdad city using nuclear track detector CR-30 (CR-39)*. Iraqi Journal of Physics المجلة العراقية الفيزياء, 2013. **11**(21): p. 54-58.
93. Jallad, K.N., *Radiation hazard indices and excess lifetime cancer risk in sand from the northern and eastern regions of Kuwait*. Environmental Earth Sciences, 2016. **75**(2): p. 1-10.
94. Karim, A.A.A.A.H.M.M.S., *Measurement of Uranium Concentrations in Human Blood in Some the Regions of Baghdad Governorate*. Ibn Al-Haitham Journal For Pure And Applied Science مجلة التطبيقية الصرفة للعلوم الهيثم ابن مجلة, 2010. **23**(2): p. 25-32.
95. Kay, D.A., *Iraq beyond the crisis du jour*. Washington Quarterly, 1998. **21**(3): p. 10-14.
96. Lerch, I.W., R.; Winick, H.; Paldy, L., *Supporting scientists and research in Iraq [2] (multiple letters)*. Science, 2003. **300**(5623): p. 1234-1235.
97. Marshall, A.C., *Analysis of Uranium Dispersal and Health Effects Using a Gulf War Case Study*. Govt Reports Announcements & Index, 2005(23): p. 212.
98. Marshall, A.C., *Gulf war depleted uranium risks*. Journal of Exposure Science and Environmental Epidemiology, 2008. **18**(1): p. 95-108.
99. Mohammed, R.S.A., R. S., *Estimation of excess lifetime cancer risk and radiation hazard indices in southern Iraq*. Environmental Earth Sciences, 2017. **76**(7).
100. Nada, F.L., AN; Enas, MY, *Uranium concentration and its associated health hazards in drinking water of Nineveh Province (Iraq)*. World Applied Sciences Journal, 2014. **31**(11): p. 1938-1944.
101. Pattison, J.E.H., R. P.; Green, S., *Enhancement of natural background gamma-radiation dose around uranium microparticles in the human body*. Journal of the Royal Society Interface, 2010. **7**(45): p. 603-611.
102. Saad M, P.K.F.W. س.ف.خ.و.ب.م.س. وارتان, *Journal of Basrah Researches (Sciences) (العلميات) البصرة ابحاث مجلة*, 2006. **32**(2B): p. 1-11.
103. Saleh, A.F.E., M. M.; Tawfiq, N. F., *Determination of uranium concentration in urine of workers in an Iraqi phosphate mine and fertilizer plants*. Journal of Radioanalytical and Nuclear Chemistry, 2013. **298**(1): p. 187-193.
104. Sallama, S.H., *Determination of Uranium Concentration In Female And Male Children's Teeth Samples Using Fission Tracks In CR-39 From Different Countries*. Journal of College of Education, 2011(5): p. 104-118.
105. Samir, M.A.-m.M.R.A.-B., *Environment Radiological Pollution from the Use of Depleted Uranium Weaponry Against Thi qar Governorate during 2003 War*. Journal of Univesity of Thi-Qar مجلة جامعة ذي العلمية قار ذي, 2006. **2**(2): p. 1-8.
106. Schott, A., et al., *Depleted Uranium (DU)—chemo-and radiotoxicity, in Uranium in the Environment*. 2006, Springer. p. 165-174.
107. Ston, R., *Nuclear control: Iraq embarks on demolition of Saddam-era nuclear labs*. Science, 2008. **321**(5886): p. 188.
108. Stone, R., *In the line of fire*. Science, 2005. **309**(5744): p. 2156-2159.
109. Szrom, F.F., G. A.; Lodde, G. M.; Parkhurst, M. A.; Daxon, E. G., *Inhalation and ingestion intakes with associated dose estimates for level II and level III personnel using capstone study data*. Health Physics, 2009. **96**(3): p. 363-379.

110. Tawfiq, N.F., *Uranium and radon concentration in ground water in Aucashat city (Iraq) and the associated health effects*. Adv Appl Sci Res, 2013. **4**(3): p. 167-171.
111. Tawfiq, N.H., NA; Bedin, S, *Determine the concentration of uranium in samples of soil and water of middle and south of Iraq using CR-39 track detector*. Journal of Baghdad for Science, 2011. **8**(2): p. 451-455.
112. Tawfiq, N.F.A., Lamya T; Al-Jobouri, Hussain A, *Uranium concentration measurements in human blood for some governorates in Iraq using CR-39 track detector*. Journal of radioanalytical and nuclear chemistry, 2013. **295**(1): p. 671-674.
113. Trešl, I.D.W., G.; Quételet, C. R.; Petrov, I.; Vanhaecke, F.; Moens, L.; Taylor, P. D. P., *Validated Measurements of the Uranium Isotopic Signature in Human Urine Samples Using Magnetic Sector-Field Inductively Coupled Plasma Mass Spectrometry*. Environmental Science and Technology, 2004. **38**(2): p. 581-586.
114. علي حسين, ا.,. Journals geographic مجلة البحوث الجغرافية, 2013. **1**(18): p. 381-413.
115. عودة سماح, ح.. Al-Mustansiriyah Journal of Science, 2012. **23**(2): p. 51-60.
116. يونس, حيدر, جعفر, موسى حسنين, وحيد, ق.ف.م, جاسم محمد, Journal of the college of basic education, الاساسية التربوية كلية مجلة, 2015. **21**(89/علمي): p. 217-224.
117. هاشم نبيل, ا., *RADIOLOGICAL AND CHEMICAL HAZARDS OF PUBLIC EXPOSURE TO URANIUM-235 AT AL-TWAITHA NUCLEAR RESEARCH SITE*. Iraqi Journal of Biotechnology المجلة العراقية للتقانات الحياتية, 2010. **9**(2): p. 226-238.
118. Alkinani, M. and B. Merkel, *Geochemistry of sediments of the Al-Batin alluvial fan, Southern Iraq*. Environmental Earth Sciences, 2018. **77**(7).
119. Almayahi, B., J.I. Hakeem, and L. Saheb, *The impact of low-level exposure to radiation in natural ecosystems of Najaf and Dhi Qar Cities, Iraq*. Iranian Journal of Medical Physics, 2018. **15**(1): p. 1-5.
120. Mohsen, A., A. Al-Khayyat, and A. Salman, *Risk of an Excess Cancer Fatality Due to Ingestion of Uranium from Some Pharmaceuticals in Iraq*. Prensa Med Argent, 2020. **106**: p. 1.
121. Najam, L.A., E.J. Mohammed, and A.S. Hameed, *Estimation of Radon Exhalation Rate, Radium Activity and Uranium Concentration in Biscuit Samples in Iraq*. Iranian Journal of Medical Physics, 2019. **16**(2): p. 152-157.

Exclude – Population

122. *CIRRPC 10th Anniversary Report. Committee on Interagency Radiation Research and Policy Coordination*. Govt Reports Announcements & Index, 1994(08): p. 54.
123. *Troops to be screened for effects of war in Iraq*. Occupational Health, 2004. **56**(3): p. 4.
124. Bollyn, C., *Depleted Uranium Blamed for Cancer Clusters Among Iraq War Vets*. Nuclear Age Peace Foundation, 2004.
125. Brower, V., *The Gulf war syndrome*. Biofutur, 1997(164): p. 25-27.
126. Clauw, D., *The health consequences of the first Gulf war*. British Medical Journal, 2003. **327**(7428): p. 1357-1358.
127. Green, G.M.G., L.; Bingham, E.; Eschenbacher, W.; Gorman, D. W.; Marcus, M.; Riley, L. W.; Schaumburg, H. H.; Singer, M. T.; Spengler, J. D.; Sutula, T. P.; Taylor, R. E.; Bascom, R.; Berg, S. W.; Blanck, R. R.; Bolton, H. T.; Cortiveau, J.;

- Daxon, E. G.; Hall, W. H., *The Persian Gulf experience and health*. Journal of the American Medical Association, 1994. **272**(5): p. 391-396.
128. Greenberg, N., *Gulf War illnesses ... the story is still alive and well*. Occupational and Environmental Medicine, 2005. **62**(3): p. 142-143.
129. Hyams, K.C., *Commentary: Adding to our comprehension of Gulf War health questions*. International Journal of Epidemiology, 2005. **34**(4): p. 808-809.
130. Knight, J., *US army survey targets Gulf War syndrome*. Nature, 2003. **421**(6922): p. 463.
131. Korenyi-Both, A.L.K.-B., A. L.; Juncer, D. J., *Al Eskan disease: Persian Gulf syndrome*. Mil Med, 1997. **162**(1): p. 1-13.
132. Lang, S., *From gulf war syndrome to balkan war syndrome*. Croatian Medical Journal, 2001. **42**(2): p. 205-209.
133. McClain, D.E., *Depleted uranium: A radiochemical toxicant?* Military Medicine, 2002. **167**(2 SUPPL.): p. 125-126.
134. McDiarmid, M.A., *Depleted uranium and public health: Fifty years' study of occupational exposure provides little evidence of cancer*. British Medical Journal, 2001. **322**(7279): p. 123-124.
135. McDiarmid, M.A.S., K. S., *Depleted Uranium (DU) Follow-up Program Update*. Govt Reports Announcements & Index, 2010(16): p. 32.
136. Mishra, R., *Editorial: Patents on decontamination of heavy metals from environment*. Recent Patents on Biotechnology, 2017. **11**(3): p. 154.
137. Nicolson, G.L.B., P.; Nasralla, M. Y.; Haier, J.; Nicolson, N. L.; Nass, M., *Gulf war illnesses: Chemical, biological and radiological exposures resulting in chronic fatiguing illnesses can be identified and treated*. Journal of Chronic Fatigue Syndrome, 2003. **11**(1): p. 135-154.
138. Nicolson, G.L.H., E.; Korenyi-Both, A.; Lopez, D. A.; Nicolson, N.; Rea, W.; Urnovitz, H., *Progress on persian gulf war illnesses reality and hypotheses*. International Journal of Occupational Medicine and Toxicology, 1995. **4**(3): p. 365-370.
139. Prophet, S., *What is Gulf War syndrome? (Part II)*. Journal of the American Health Information Management Association, 1998. **69**(8): p. 77-86.
140. Sartin, J.S., *Gulf War Syndrome: The final chapter?* Mayo Clinic Proceedings, 2006. **81**(11): p. 1425-1426.
141. Squibb, K.S.O., M.; Gucer, P.; Engelhardt, S.; McDiarmid, M., *Utility Of Patient Mailed Urine Specimens For Renal Biomarker Analysis In Depleted Uranium (DU) Exposed 1991 Gulf War Veterans*. Toxicol Sci, 2005. **84**(1-S): p. 35.
142. Ushakov, I.B.A.e., R. V.; Berezin, G. I.; Zuev, V. G., *Depleted uranium: radiation and ecological safety aspects*. Voenno-meditsinskii zhurnal, 2003. **324**(4): p. 56-58, 80.
143. Vastag, B., *Battlefield Uranium Doubts Linger for Troops*. Journal of the American Medical Association, 2003. **289**(13): p. 1621-1623.
144. Vasterling, J.J.B., J. D., *The impact of the 1991 Gulf War on the mind and brain: Findings from neuropsychological and neuroimaging research*. Philosophical Transactions of the Royal Society B: Biological Sciences, 2006. **361**(1468): p. 593-604.

145. Wadman, M., *US panel draws blank on Gulf War symptoms*. *Nature*, 2000. **407**(6801): p. 121.
146. Young, R.C.J.R., R. E.; Hodge, W. D., *Threats to respiratory health in the persian gulf region*. 85th Annual Scientific Assembly of the Southern Medical Association in Conjunction with the Medical Association of Georgia, Atlanta, Georgia, USA, November 16-19, 1991. *South Med J*, 1991. **84**(9): p. 2S12.
147. Zunic, S., *Lupus erythematosus cell phenomenon in pediatric bronchoalveolar lavages: Possible manifestation of early radioadaptive response in radiation-induced alveolitis*. *Journal of Biological Regulators and Homeostatic Agents*, 2013. **27**(2): p. 389-398.

Exclude – Study Design: Primary

148. *UNEP Depleted Uranium Awareness*. United Nations Environment Programme.
149. *Mixed messages about depleted uranium*. *Lancet Oncol*, 2001. **2**(2): p. 65.
150. *Depleted Uranium And use an American in Iraq*. *Journal of Political Sciences مجلة السياسية العلوم*, 2014(48): p. 65-85.
151. Abbott, A., *WHO plans study of Gulf War fallout*. *Nature*, 2001. **413**(6852): p. 97.
152. Airhart, M., *Cleaning Up after War*. *Scientific American*, 2003. **289**(4): p. 44-45.
153. Aitken, M., *Gulf war leaves legacy of cancer*. *Bmj*, 1999. **319**(7207): p. 401.
154. Åkerblom, G. *Depleted uranium - Experience of the United Nations environmental programme missions*. in *8th International Symposium on the Natural Radiation Environment, NRE VIII*. 2008. Buzios, Rio de Janeiro.
155. Al Ani, A.-H.B., Joanne, *Uranium in Iraq. The poisonous legacy of the Iraq Wars*. 2009: Florida: Vandenplas Publishing.
156. Al-Fahaad, H.A., *Depleted uranium. Is it potentially involved in the recent upsurge of malignancies in populations exposed to war dust?* *Saudi Medical Journal*, 2012. **33**(9): p. 1028.
157. Al-Mendalawi, M.D., *Epidemiology of neural tube defects*. *Saudi Medical Journal*, 2015. **36**(3).
158. Bertell, R., *Depleted uranium: All the questions about DU and Gulf War syndrome are not yet answered*. *International Journal of Health Services*, 2006. **36**(3): p. 503-520.
159. Birchard, K., *Does Iraq's depleted uranium pose a health risk?* *Lancet*, 1998. **351**(9103): p. 657.
160. Brown, V.J., *Reconstructing the environment in Iraq*. *Environmental Health Perspectives*, 2004. **112**(8): p. A464.
161. Brumfiel, G., *Iraqi looters spark alert over radiation risks*. *Nature*, 2003. **423**(6938): p. 370.
162. Chesser, R.K.R., B. E.; Bondarkov, M.; Shubber, E.; Phillips, C. J., *Piecing together Iraq's nuclear legacy*. *Bulletin of the Atomic Scientists*, 2009. **65**(3): p. 19-33.
163. Collier, R., *Iraq Links Cancers to Uranium Weapons*. *San Francisco Chronicle*, 2003.
164. Edwards, R., *WHO 'suppressed' scientific study into depleted uranium cancer fears in Iraq*. *Sunday Herald*. <http://www.sundayherald.com/40096>, 2004.

165. Fahey, D., *The Use of Depleted Uranium in the 2003 Iraq War: An Initial Assessment of Information and Policies*. Berkeley, Calif., June, 2003. **24**.
166. Fasy, T. *The recent epidemic of pediatric malignancies and congenital malformations in Iraq: The biological plausibility of depleted uranium as a carcinogen and a teratogen*. in *Iraqi-American Academics' Symposium for Peace*. 2003.
167. Graham-Rowe, D., *Depleted uranium casts a shadow over peace in Iraq*. New Scientist, 2003. **178**(2391): p. 4-4.
168. Hall, S.O., D., *Killing them softly*. Nursing times, 1999. **95**(34): p. 18.
169. Hameed Hamed Al-Sa'adoon محمد حميد ا., *U.S.A. use of depleted uranium in Iraq*. Journal of International studies مجلة دولية دراسات / مركز عن تصدر الأستراتيجية الدراسات مركز عن تصدر / الدولية 2012(54): p. 83-100.
170. Hirschfield, R., *An Arab-American priest, depleted uranium, and Iraq*. The Washington Report on Middle East Affairs, 2005. **24**(8): p. 29.
171. Huggan, P., *Health effects of war are devastating: Pre-emptive conflict demands pre-emptive criticism by the medical profession [4]*. Internal Medicine Journal, 2007. **37**(12): p. 839-840.
172. Kapp, C., *WHO sends team to Iraq to investigate effects of depleted uranium*. Lancet., 2001. **358**(9283): p. 737. [Lancet (London, England)].
173. Kathren, R.L., *Depleted uranium: A problem of perception rather than reality*. Radiation Protection Dosimetry, 2001. **95**(1): p. 3-4.
174. Kazashi, N., *The invisible 'internal radiation' and the nuclear system: Hiroshima-Iraq-Fukushima*. Ethics, Policy and Environment, 2012. **15**(1): p. 37-43.
175. Kincaid, R., *The devastating impact of depleted uranium weapons on civilian and military personnel: After the war, does peace have to be hell, too?* International Journal of Interdisciplinary Social Sciences, 2008. **3**(8): p. 49-58.
176. Levy, B.S.S., V. W., *Adverse health consequences of the Iraq War*. The Lancet, 2013. **381**(9870): p. 949-958.
177. Mathews, J., *Radioactive Bullets Raise Cancer Fears*. Journal of the National Cancer Institute, 1993. **85**(13): p. 1029-1030.
178. Menkhi, S.A.S., F. H.; Almayahi, B. A., *Radiation pollution and cancer risks in Sulaimaniyah and Ninawa Cities, Iraq*. Annual Research and Review in Biology, 2017. **18**(4).
179. Mir, L., *Risks related to the military use of depleted uranium during the Gulf war*. Environnement, Risques et Sante, 2008. **7**(4): p. 241-242.
180. Moret, L., *From Hiroshima to Iraq, 61 years of uranium wars*. A suicidal, genocidal, omnicidal course, Global Research, 2007.
181. Moszynski, P., *Royal Society warns of risks from depleted uranium*. BMJ (Clinical research ed.), 2003. **326**(7396): p. 952.
182. Moszynski, P., *Deadly radiation ignored in Iraq*. BMJ (Clinical research ed.), 2003. **327**(7405): p. 11.
183. Mould, R., *Depleted uranium and radiation-induced lung cancer and leukaemia*. The British journal of radiology, 2001. **74**(884): p. 677-683.
184. Natali, L., *Contemporary wars and environmental consequences. A green criminological approach*. Rassegna Italiana di Criminologia, 2016. **10**(3): p. 209-218.

185. Nau, J.Y., *The Gulf War syndrome does exist*. *Revue Medicale Suisse*, 2008. **4**(184): p. 2764.
186. Peterson, S., *The Gulf war battlefield still "hot" with depleted uranium*. *Middle East Report*, 1999(211): p. 2.
187. Pivovarov Iu, P.A.-S., A. A.; Sheina, N. I., [*Problem of unpredictable anthropogenic exposure to the state of natural environment in countries of southwest Asia*]. *Gig Sanit*, 2013(6): p. 21-5.
188. Rokke, D., *Depleted uranium and its effects in Iraq*. *Proceedings, The Campaign Against Sanctions on Iraq*, 1999: p. 121-143.
189. Roth, P., *Uranium exposure and Golf War Syndrome*. *Gynakologische Praxis*, 2005. **29**(1): p. 197-198.
190. Shanoon, F.H.M., S. A.; Almayahi, B. A.; Dawood, A. S., *Spatial and temporal variability of environmental radioactivity in basra and baghdad cities, iraq*. *Annual Research and Review in Biology*, 2017. **21**(6).
191. Shelleh, H.H., *Depleted Uranium is it potentially involved in the recent upsurge of malignancies in populations exposed to war dust?* *Saudi Medical Journal*, 2012. **33**(5): p. 483-488.
192. Spratt, B.G., *Invited Editorial: Depleted uranium munitions - Where are we now?* *Journal of Radiological Protection*, 2002. **22**(2): p. 125-129.
193. Van Hoof, R., *Gulf syndrome...Balkans syndrome...chronic fatigue syndrome*. *Bulletin et mémoires de l'Académie royale de médecine de Belgique*, 2000. **155**(7-9): p. 281-291; discussion 291-294.
194. Wagner, F.T., V, *The doctor, the depleted uranium and the dying children, documentary film pro-duced for German television and released by Ochoa-Wagner production in 2004 in Germany, exposes the use and impact of radioactive weapons during the current war against Iraq*. 2005.
195. Wakeford, R., *Depleted uranium*. *Journal of radiological protection : official journal of the Society for Radiological Protection*, 2001. **21**(1): p. 76-77.
196. Walker, M., *Truth and technology at war*. *New Scientist*, 2003. **180**(2426-2428): p. 12-13.
197. Webster, P.C., *Questions raised over Iraq congenital birth defects study*. *The Lancet*, 2013. **382**(9899): p. 1165-1166.
198. Williams, D., *Hazards of Uranium weapons in proposed war on Iraq*. sept. 22nd, 2002.
199. Zolotov, Y.A., *Mass spectrometry, enrichment of uranium, and Iraq*. *Journal of Analytical Chemistry*, 2008. **63**(9): p. 823.
200. Zwijnenburg, W., *In a State of Uncertainty: Impact and Implications of the Use of Depleted Uranium in Iraq*. 2013: IKV Pax Christi.
201. الفلاحي, M.M.E., م.م.ا. ف. و. ف. أ. ع. أ. أ. O.A.A., *DEPLETED URANIUM AND BIOTA BIOLOGY*. *The Iraqi Journal of Agricultural Science مجلة العلوم الزراعية العراقية*, 2010. **41**(1): p. 1-15.
202. *Worldwide Emerging Environmental Issues Affecting the U.S. Military. July 2005 Report*. *Govt Reports Announcements & Index*, 2005(02): p. 18.
203. Koutras, A., et al., *Changes in the incidence of breast cancer due to the use of radioactive materials for warfare or nuclear and environmental accidents over the last 60 years in Europe and Asia*. *Journal of B.U.ON.*, 2019. **24**(1): p. 5-10.

Exclude – Study Design: Comparator

204. Al-Abbasi, D.S.A.-J., A. A.; Al-Toriahi, K. M.; Jabor, T. A.; Yasseen, A. A., *Expression of VEGF in urinary bladder transitional cell carcinoma in an Iraqi population subjected to depleted uranium: An immunohistochemical study*. Applied Immunohistochemistry and Molecular Morphology, 2009. **17**(4): p. 307-3112.
205. Al-Dujaily, E.A.A.-J., A. A.; Pierscionek, T.; Yasseen, A. A., *High prevalence of HER-2/neu overexpression in female breast cancer among an Iraqi population exposed to depleted uranium*. J Carcinog, 2008. **7**: p. 8.
206. Al-Hadithi, T.S.A.-D., J. K.; Saleh, A. M.; Shabila, N. P., *Birth defects in Iraq and the plausibility of environmental exposure: A review*. Conflict and Health, 2012. **6**(1).
207. Ali Jabir Neima, A.-k.L., H. Al-Kelabi; Esraa, A. A. L. Dujaily, *HER-2/neu overexpression in correlation to Vascular Endothelial Growth Factor ,grade and stage of Non other wised specified Invasive ductal carcinoma*. Al-Qadisiah Medical Journal *مجلة الطبية القادسية*, 2010. **6**(10): p. 27-37.
208. Ali, L.H.A., H. L., *Histological alterations in leukocytes of patients suffering leukemia by scanning electron microscope*. Journal of Pharmaceutical Sciences and Research, 2017. **9**(10): p. 1698-1700.
209. Al-Mumen, M.M.A.-J., A. A.; Jumaa, A. S.; Al-Toriahi, K. M.; Yasseen, A. A., *Exposure to depleted uranium does not alter the co-expression of HER-2/neu and p53 in breast cancer patients*. BMC Research Notes, 2011. **4**.
210. Bell, D.E., *Public health controversy of radioactive warfare: Depleted uranium and displaced discourse in Medically Unexplained Chronic Syndromes (MUCS)*. 2013, State University of New York at Buffalo: Ann Arbor. p. 478.
211. FATHI, R.A.G., DOUG; AL-SALIH, HANA, *The accumulation of uranium in wild plants (Lagonychium farctum) in Nineveh Governorate/Iraq*.
212. Fathi, R.A.M., L. Y.; Al-Salih, H. S.; Godbold, D., *Environmental pollution by depleted uranium in Iraq with special reference to Mosul and possible effects on cancer and birth defect rates*. Medicine, Conflict and Survival, 2013. **29**(1): p. 7-25.
213. Hade, H.A.J., R. H.; Hatrosh, S. J., *CLEC4E as novel tumor marker. A biochemical study for prediction acute lymphocytic leukemia at iraqi children*. Journal of Pharmaceutical Sciences and Research, 2018. **10**(3): p. 556-561.
214. Ifad Kerim, A.-s.M.A., Muhsin; Mohammed Sabri, A. Razak, *Immunological Study On Breast Cancer In Hilla Province*. Kerbala Jorunal of Medicine *مجلة كربلاء الطبية*, 2011. **4 no 2**(10): p. 1140-1145.
215. Khudheir, M.Y., Ahmad Shukri; Ahm, Fauziah, *Hazard Assessment of Humanity as result of Biological Contamination with Uranium at Iraq Southern*. International Journal of Biological Engineering, 2013. **3**(2): p. 18-25.
216. Mahmoud, A.A. and I.M. Abdallah, *Effect of Depleted Uranium As One of Nuclear Waste on The Collapse of The Immune System and The Spread of Cancerous Diseases and the impact of higher cost through treatment*. Scientific Journal of Karbala University, 2012. **10**(2): p. 88-101.

217. Marouf, B. *Environmental Impact of Depleted Uranium (DU) Contamination in Iraq*. in *The Conference on The Effects of The Use of Depleted Uranium Weaponry on Human and Environmental in Iraq*. 2000.
218. Nsireen Hadi Rasheed رشيد هادي نسرین, *The spatial Distribution for the Radiation pollution in Iraq*. Journal of Research Diyala humanity مجلة الانسانية للبحوث ديالى, 2013(57): p. 729-763.
219. Salih, N.F.J., M. S., *Novel method to assessing and the impact of alpha emitter's concentration of the uterus on women fertility in Iraqi Kurdistan region*. Journal of Radioanalytical and Nuclear Chemistry, 2013. **298**(2): p. 755-761.
220. Salih, N.F.J., M. S., *Investigation of alpha emitters in fresh and powdered blood of fertile women: An in vitro application of CR-39 NTDs*. Journal of Radioanalytical and Nuclear Chemistry, 2014. **300**(2): p. 693-699.
221. Samir, M.K.A., F. Shehab; Akeel, H. A. Al-Asse, *A molecular Study of Azoospermia & Sever Oligospermia in a sample of infertile Males Al-Anbar Governorate*. Engineering and Technology Journal مجلة الهندسة والتكنولوجيا, 2012. **30**(12): p. 2116-2124.
222. ك, سالم محمود (C R- 3 9). Journal of College of Education, 2010(3): p. 161-170.
223. Al-Hamzawi, A.A.J., M. S.; Tawfiq, N. F., *Uranium concentration in blood samples of Southern Iraqi leukemia patients using CR-39 track detector*. Journal of Radioanalytical and Nuclear Chemistry, 2014. **299**(3): p. 1267-1272.
224. Al-Kzayer, L.a.F.Y., et al., *Asymmetrically enlarged parietal foramina in a rare case of Goldenhar syndrome with a possible etiopathogenesis*. Oncotarget, 2018. 9(2): p. 2962-2968.
225. Mahmood, A.S., O.S. Shafeq, and M.S. Shafiq, *Uranium concentration variation dependency on gender correlated with age of bladder cancer patient*. International Journal of Research in Pharmaceutical Sciences, 2019. 10(3): p. 1730-1734.