Total Diet Approach to Cancer Prevention: A Narrative Review of the Literature Since 2011 Concerning the Association of Colorectal Cancer and Meat Consumption

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Abstract
Diet serves as a primary prevention approach to reduce the global burden of cancer. In 2007, the World Cancer Research Fund and the American Institute for Cancer Research published the Second Expert Report (SER) Food, Nutrition, Physical Activity, and the Prevention of Cancer: A Global Perspective outlining lifestyle recommendations for primary cancer prevention. Results support a strong link between red and processed meat and colorectal carcinogenesis. Findings from the Colorectal Cancer 2011 Report: Food, Nutrition, Physical Activity, and the Prevention of Colorectal Cancer, a review conducted through the WCRF Continuous Update Project (CUP), strengthened the evidence and supported the conclusions found in the SER. This review explored the available evidence since the publication of the 2011 CUP report and provides an update of the literature, specific to colorectal cancer (CRC) and diet. Furthermore, several proposed mechanisms, including heterocyclic amines (HCAs), polyaromatic hydrocarbons (PAHs), N-nitroso compounds (NOCs), and heme iron, may explain the effects of meat on the cancer process. The studies reviewed continue to support the causal link between red and processed meat consumption and CRC. The most recent literature supports the preventative role of consuming a plant-based diet low in red and processed meat for overall cancer prevention.

Keywords: Colorectal cancer; Cancer prevention; Diet; Iron; Cancer risk


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Introduction
Although the exact causes of cancer are unknown and genetic predisposition may afford an elevated risk, approximately 30-40 percent of cancers are preventable over time through lifestyle behaviors, such as smoking, diet, and physical activity [1]. Specifically, energy dense dietary consumption and physical inactivity lead to weight gain. Being overweight and obese attributes to the development of one-third of cancer cases worldwide [1]. Previous studies show that obesity is strongly associated with the development of breast, prostate, endometrial, esophageal, and pancreatic cancer. According to the National Cancer Institute (NCI), approximately 85,000 new cancer cases were attributed to obesity in 2007 in the US, with a
projection of obesity leading to about 500,000 additional cancer cases by 2030 [2]. Therefore, maintaining a normal weight and preventing weight gain over the lifespan through healthy eating and daily physical activity reflect foundational evidence-based strategies for cancer prevention.

In 2007, the World Cancer Research Fund (WCRF) and the American Institute for Cancer Research (AICR) published the Second Expert Report (SER) Food, Nutrition, Physical Activity, and the Prevention of Cancer: A Global Perspective, which reflects a synthesis of the current literature and outlines the top recommendations for cancer prevention. In general, the SER committee recommended following a plant-based diet by eating a variety of vegetables, fruits, whole grains, and legumes, while limiting the consumption of red meat1 and avoiding processed meat2. Some of the strongest evidence from this report reveals a significant ‘cause and effect’ relationship between the consumption of red and/or processed meat consumption and the development of colorectal cancer (CRC) [1]. The American Cancer Society (ACS) projected that in 2015, 69,090 men and 63,610 women will receive a CRC diagnosis and ultimately close to 50,000 combined deaths from CRC will occur [3]. As such, it is imperative to identify prevention strategies to decrease the overall burden of CRC over time. Through the Continuous Update Project (CUP), the WCRF went on to publish the 2011 Food, Nutrition, Physical Activity, and the Prevention of Colorectal Cancer, which includes new scientific findings since the publication of the original report. Consistent with findings from the SER, the CUP panel also concluded that both red and processed meat continue to be convincing causes of CRC [4]. New research has emerged since the publication of this report regarding the associations between diet and CRC. Therefore, the objective of this narrative review is to provide a review of the literature since the 2011 CUP report, including the proposed mechanisms and biological plausibility of meat consumption and cancer development.

Methodology

A comprehensive literature search was conducted using the PubMed (NIH) database (2009-2015). The search was limited to human studies published in English. Keyword combinations of the medical subject headings (MeSH) included: "processed meat", "red meat", "red meat intake", "processed meat intake", "frequency of meat consumption", "cancer", "colorectal cancer", "colorectal cancer risk", "heterocyclic amines", "HCA", "well-done meat", "cooked meat", "cooking methods" "cooking practices", "iron", "heme iron", "nitrate", "nitrite", or "N-nitroso compounds". A secondary search was conducted by reviewing the references of articles to identify further manuscripts for inclusion and critical review.

1According to the SER ‘red meat’ refers to “beef, pork, lamb, and goat from domesticated animals including that contained in processed foods” [1].

2According to the SER ‘processed meat’ refers to “meat preserved by smoking, curing or salting, or addition of chemical preservatives, including that contained in processed foods” [1].

Results

Figure 1 depicts the search strategy and paper selection process. Initially, 858 papers were reviewed; however, 20 papers were deemed evaluable, after duplicative papers and further search restrictions were applied. The following sections summarize the results of the literature review including compounds formed from high-heat cooking methods, nitrates/nitrites in processed meat, and heme iron. Tables 1-4 provide a summary of the studies, describing the study design, key characteristics of the population, dietary assessment methodologies, results and significant findings.

Meat consumption and colorectal cancer

The negative effects of animal-based protein are most pronounced in CRC [1]. The SER states that there is convincing evidence to support red meat as a cause of CRC, with cohort data demonstrating a dose-response relationship. The WCRF/AICR guideline recommends limiting consumption of red meat, such as beef, pork, and lamb, to no more than 18 ounces per week and avoiding processed meats, such as ham, bacon, salami, hot dogs, and sausages [1]. Since the publication of the SER, six studies have been published, four cohort studies and two case-control studies on the link between CRC and red meat consumption. Of these studies, five supported that consuming red or processed meat was associated with development of CRC, while one case control investigation concluded that there was insufficient evidence to support this relationship. These studies are summarized in Table 1.

The five studies in support of the relationship between meat and elevated CRC risk included three cohort and two case-control studies. In a prospective cohort investigation, the role of meat consumption and colorectal cancer...
consumption and risk of adenoma in the distal colon and rectum was investigated using 1008 individuals from the Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial with incident distal colorectal adenoma [5]. Positive associations were observed for red (odds ratio (OR) 1.22, 95% CI 0.98-1.52) and processed meat (OR 1.23, 95% CI 0.99-1.54), as well as heme iron (OR 1.23, 95% CI 0.99-1.52) and nitrate/nitrite (OR 1.22, 95% CI 0.94-1.53). The findings indicate that several meat-related components may be most relevant to early neoplasia in the rectum.

### Table 1 Association between red, processed meat consumption and CRC.

<table>
<thead>
<tr>
<th>Citation/Topic</th>
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<th>Dietary Assessment Method</th>
<th>Results</th>
<th>Major Findings/Conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ferrucci et al. [5]</td>
<td>Prospective cohort study of participants in the screening arm of the Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial who underwent baseline and follow-up sigmoidoscopy 1008 w/ incident distal CR adenoma</td>
<td>137-item FFQ on usual intake of foods and beverages during the past year</td>
<td>Positive associations found for red (OR 1.22, 95% CI 0.98-1.52) and processed meat (OR 1.23, 95% CI 0.99-1.54), as well as heme iron (OR 1.23, 95% CI 0.99-1.52) and nitrate/nitrite (OR 1.22, 95% CI 0.94-1.53)</td>
<td>Findings indicate that several meat-related components may be most relevant to early neoplasia in the rectum</td>
</tr>
<tr>
<td>Takachi et al. [6]</td>
<td>Prospective cohort study; Japanese cohort of 80,658 men and women age 45-74</td>
<td>138-item FFQ, including 16 meat items</td>
<td>Significant association found between higher red meat consumption among women (HR 1.48, 95% CI 1.01-2.17) and higher total meat consumption among men (HR 1.44, 95% CI 1.06-1.98)</td>
<td>Red meat may increase the risk of colon cancer in middle-aged Japanese. The highest quintile of red meat consumption may be considered moderate by Western standards</td>
</tr>
<tr>
<td>Bernstein et al. [7]</td>
<td>Prospective cohort study of participants in the Nurses’ Health Study (NHS) (n=87,108 women 1980-2010) and the Health Professionals Follow Up Study (HPFS) (n=47,389 men 1986-2010) Combined cohorts – 2,731 CRC cases</td>
<td>NHS: validated 61-item FFQ in 1980, expanded FFQ every 4 years from 1984-2010 to update dietary intake HPFS: validated 131-item FFQ, administered every 4 years from 1990-2010</td>
<td>Multivariable analysis showed modest significant association found between processed red meat and CRC (HR 1.15, 95% CI 1.01-1.32)</td>
<td>A significant and positive association between processed red meat and CRC, particularly distal colon cancer, was observed</td>
</tr>
<tr>
<td>Aune et al. [8]</td>
<td>Multisite case-control study of 11 cancer sites in Uruguay (1996-2004); 3539 cancer cases and 2032 hospital controls</td>
<td>64-item FFQ covering dietary intake one year before diagnosis</td>
<td>High intake of red and processed meat was significantly associated (OR 3.83, 95% CI 2.37-6.20; OR 2.15, 95% CI 1.49-3.11 respectively) with an increase in CRC</td>
<td>Results confirm the association between red and processed meat consumption and CRC risk, as well as several other cancer sites – oral cavity and pharynx, esophagus, larynx, stomach, lung, breast, prostate, bladder, and kidney</td>
</tr>
<tr>
<td>De Stefani et al. [9]</td>
<td>Case-control study in Uruguay: 611 CRC cases and 1,362 controls</td>
<td>64-item FFQ</td>
<td>The highest risk was positively associated with meat-based pattern (OR 1.63, 95% CI 1.22-2.18), where the plant-based pattern (OR 0.60, 95% CI 0.45-0.81) was strongly protective with CRC risk</td>
<td>Meat-based pattern suggestive of highest CRC risk; since HCAs are formed in well-done meat, it is suggestive that meat-based pattern could be an important etiologic agent for CRC</td>
</tr>
<tr>
<td>Spencer et al. [10]</td>
<td>Case-control study with 579 cases of CRC matched with 1996 controls</td>
<td>4-7 day food diaries–disaggregated weights of meat, poultry, and fish from composite foods to investigate dose-response relationships</td>
<td>Disaggregated intakes were moderately low (e.g. mean red meat intakes were 38.2 g/day among male and 28.7 g/day among female controls) little evidence of association between red (OR 0.88, 95% CI 0.68-1.15) and processed meat (OR 0.97, 95% CI 0.84-1.12) and risk of CRC</td>
<td>This study using pooled data from prospective food diaries, among cohorts with low to moderate meat intakes, shows little evidence of association between consumption of red and processed meat and CRC risk</td>
</tr>
</tbody>
</table>
and the findings supported a significant association between higher red meat consumption among women (HR 1.48, 95% CI 1.01-2.17) and higher total meat consumption among men (HR 1.44, 95% CI 1.06-1.98) [6]. The increased risk of colon cancer with meat consumption was found in this population who have relatively low intakes of red meat compared to Western standards. Additionally, in a study of participants from the Nurses’ Health Study and the Health Professionals Follow Up Study, multivariable analysis of a combined cohort of 2,731 CRC cases found a modest significant association between processed red meat and CRC (HR 1.15, 1.01-1.32) [7].

Two case-control studies were conducted in Uruguay. Aune et al. [8] assessed the associations between cancer risk and meat intake including 11 cancer sites and comprising 3,539 cancer cases and 2,032 hospital controls in Uruguay between 1996 and 2004. High intake of red and processed meat was significantly associated (OR 3.83, 95% CI 2.37-6.20; OR 2.15, 95% CI 1.49-3.11 respectively) with an increase in CRC. The case-control study by De Stafani et al. [9] explored the association between CRC risk and nutrient-derived dietary patterns. The study analyzed 611 cases of CRC and 1,362 controls and retained three factors: Meat-based, plant-based, and carbohydrate dietary patterns. The highest risk of CRC was positively associated with the meat-based pattern (OR 1.63, 95% CI 1.22-2.18), whereas a strong protective effect was found with the plant-based pattern (OR 0.60, 95% CI 0.45-0.81) [9]. The findings of this study suggested that a meat-based dietary pattern was an etiologic agent for CRC.

Spencer et al. [10] matched 579 cases of CRC with 1,996 controls and used food diaries to examine dietary intake over 4-7 days. The results showed limited evidence to support the association between consumption of red (OR 0.88, 95% CI 0.68-1.15) and processed meat (OR 0.97, 95% CI 0.84-1.12) and risk of CRC [10]. The authors noted that disaggregated intakes of red meat, or estimated amounts separated from mixed dishes (which tend to lead to overestimation of amounts consumed), were moderately low with mean red meat intakes of 38.2 g/day for male and 28.7 g/day for female controls [10]. Collectively, the majority of the studies included in this update since the CUP Report continue to support the cause and effect relationship between CRC risk and meat intake and the recommendation to limit red and processed meat for the prevention of CRC.

Three Underlying Mechanisms and Biological Plausibility

Heterocyclic amines/polycyclic aromatic hydrocarbons and cooking methods

Heterocyclic amines (HCAs) and polycyclic aromatic hydrocarbons (PAHs) are chemicals formed from the reaction of creatine or creatinine, amino acids, and sugar in muscle meat using high-temperature cooking methods, such as pan frying or grilling directly over an open flame. The most important factor in the formation of HCAs and PAHs is temperature [11]. These mutagenic chemicals may cause DNA damage after specific enzymes in the body metabolize them through a process called “bioactivation.” Evidence shows that HCA and PAH exposure can cause cancer in animal models [12]. While a definitive link between exposure to HCAs and PAHs has not been established in human studies, as it is difficult to determine the exact level of exposure to these chemicals, population based epidemiologic studies estimate exposure using detailed questionnaires. Table 2 summarizes the literature on six studies on the association between HCAs and CRC since the publication of the CUP Report. Of the studies assessed, three supported the association between HCAs and CRC and three indicated the evidence insufficient to support the association.

One cohort study and two case-control studies supported the association between HCAs and risk of CRC. A prospective cohort study analyzed data collected from the European Prospective Investigation into Cancer and Nutrition (EPIC) – Heidelberg Study [11]. HCA intake was estimated in 21,452 participants aged 35-65 who completed a follow-up food frequency questionnaire (FFQ) that included detailed questions on meat preparation methods and preferred degree of browning. Analysis showed that 2-amino-1-methyl-6-phenylimidazo[4,5,b]pyridine (PhIP), a HCA, was the most abundant dietary HCA associated with increased colorectal adenoma risk (RR 1.47, 95% CI 1.13-1.93), but 2-amino-3,8-dimethylimidazo[4,5-f]quinoxaline (MeIQx) (RR 1.27, 95% CI 0.97-1.68) and 2-amino-3,4,8-dimethylimidazo[4,5-f]quinoxaline (DiMeIQx) (RR 1.18, 95% CI 0.92-1.53), other HCAs, did not show statistically significant observations [11]. The results of this study support a positive association between intake of specific HCAs and risk of CRC. A case-control study assessed the associations between meat-related compounds as underlying mechanisms for CRC risk [12]. Participants included 989 cases and 1,033 healthy controls that completed a FFQ with a meat-specific module. Positive associations were observed for DiMeIQx and colorectal (OR 1.36, 95% CI 1.02-1.82), distal colon (OR 1.66, 95% CI 1.06-2.59), and rectal (OR 1.54, 95% CI 1.02-2.33) tumors [12]. In the case-control study by Helmus et al. [13], 1,062 incident colon cancer cases and 1,645 population controls completed a meat preparation questionnaire and reported statistically significant associations between dietary intake of MeIQx (adjusted odds ratio (aOR) 1.88 CI 1.45-2.43), DiMeIQx (aOR 1.73 CI 1.34-2.23), meat-derived mutagenic activity (aOR 1.84 CI 1.42-2.39) and colon cancer carcinogenesis [13].

Two cohort studies and one case-control study concluded that there was insufficient evidence to support the association between HCAs and risk of CRC. A prospective cohort study conducted within the Multiethnic Cohort Study examined the association between CRC risk and consumption of total, red, or processed meat in 165,717 participants who completed a detailed FFQ [14]. This study assessed whether a greater estimated intake of HCAs had an association with risk of CRC among 131,763 participants who completed a follow up questionnaire with an added meat-cooking module. While the relationship between meat or HCAs was not statistically significant (RR 0.90, 95% CI 0.76-1.05), the authors noted that they could not rule out the possibility that...
meat and HCAs had a modest effect on the development of CRC due to residual confounding (e.g., differences in the approaches used to classify and define red and processed meat, variable recipes, etc.) [14]. Parr et al. [15] examined associations of meat intake with incident cancer of the proximal colon, distal colon, and rectum in the population-based Norwegian Women and Cancer cohort. Cooking methods of meat were also examined in a subsample of the cohort. The findings of the study do not support an association between CRC and intake of red meat or cooking methods (HR 0.70, 95% CI 0.33-1.48) [15]. Tabatabaei et al. [16] conducted a population-based case-control study in the Western Australian Bowel Health Study that investigated the association between meat consumption, cooking practices, and CRC risk. The findings from the study do not support the hypothesis that meat intake is a risk factor for CRC (OR 0.73, 95% CI 0.53-1.01) [16].

### Nitrates/Nitrites and processed meat

Processed meats are preserved with nitrite, which can react with degradation products of amino acids to form N-nitroso compounds and CRC risk [11]. Parr et al. [15] examined associations of meat intake or processed meat intake or cooking methods (HR 0.70, 95% CI 0.33-1.48) [16]. Tabatabaei et al. [16] conducted a population-based case-control study in the Western Australian Bowel Health Study that investigated the association between meat consumption, cooking practices, and CRC risk. The findings from the study do not support the hypothesis that meat intake is a risk factor for CRC (OR 0.73, 95% CI 0.53-1.01) [16].

### Table 2: Association between heterocyclic amines/polycyclic aromatic hydrocarbons, cooking methods, and CRC.

<table>
<thead>
<tr>
<th>Author/Topic</th>
<th>Study Design/Population</th>
<th>Dietary Assessment Method</th>
<th>Results</th>
<th>Major Findings/Conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rohrmann et al.</td>
<td>Prospective cohort study; 25,540 participants of the European Prospective Investigation into Cancer and Nutrition-Heidelberg cohort study, 21,452 completed follow-up FFQ</td>
<td>FFQ on meat consumption, applied preferred degree of browning</td>
<td>Intake of PhIP associated with increased risk of CR adenoma (RR 1.47, 95% CI 1.13-1.93) but no statistically significant associations observed for MeIQx (RR 1.27, 95% CI 0.97-1.68) and DiMeIQx (RR 1.18, 95% CI 0.92-1.53)</td>
<td>Support data from case-control studies of a positive association between HCA intake and CRC adenoma risk</td>
</tr>
<tr>
<td>Miller et al.</td>
<td>Case-control; 989 cases/1,033 healthy controls</td>
<td>FFQ w/a meat-specific module</td>
<td>Significant positive association was observed for HCA DiMeIQx and colorectal (OR 1.36, 95% CI 1.02-1.82), distal colon (OR 1.66, 95% CI 1.06-2.59), and rectal (OR 1.54, 95% CI 1.02-2.33) tumors</td>
<td>HCAs, PAHs, nitrites, and nitrates may be involved in CRC etiology</td>
</tr>
<tr>
<td>Helmus et al.</td>
<td>Case-control study of 1,062 incident colon cancer cases and 1,645 population controls</td>
<td>175-item FFQ and validated meat preparation questionnaire adapted from a National Cancer Institute HCA concentration database</td>
<td>Dietary intake of MeIQx (adjusted odds ratio (aOR) 1.88, 95% CI 1.45-2.43), DiMeIQx (aOR 1.73 95% CI 1.34-2.23), and meat-derived mutagenic activity (aOR 1.84 95% CI 1.42-2.39) showed statistically significant associations with risk of colon cancer</td>
<td>Results supports that HCAs and PAHs derived from red meat, but not white meat, are a potential environmental pathway of colon cancer carcinogenesis</td>
</tr>
<tr>
<td>Ollberding et al.</td>
<td>Prospective cohort study conducted within the Multiethnic Cohort Study; 165,717 participants–greater consumption of total, red, or processed meat associated with CRC risk and 131,763 participants who completed a follow-up questionnaire that included a meat-cooking module; 3,404 and 1,757 CRC cases</td>
<td>Detailed quantitative FFQ that obtained the frequency and quantity of food items consumed during the preceding year</td>
<td>No association with CRC risk detected for density-adjusted total meat or processed meat intake or for total or specific HCA intake (RR 0.90, 95% CI 0.76-1.05) comparing quintiles of dietary exposure or using continuous variables</td>
<td>Results do no support a role for meat or HCAs from meat in etiology of CRC; cannot rule out modest effect</td>
</tr>
<tr>
<td>Parr et al.</td>
<td>Prospective cohort study; population-based Norwegian Women and Cancer cohort; 84,538 women; 459 colon and 215 rectal cancer cases with follow-up</td>
<td>FFQ that covered habitual, but not total diet, during the previous year</td>
<td>The findings of the study do not support an association between the risk of CRC and intake of red meat or cooking methods (HR 0.70, 95% CI 0.33-1.48)</td>
<td>Did not support an association between CRC risk and intake of red meat or meat cooking methods, but a high processed meat intake was associated with increased risk of proximal colon, distal and rectal cancer</td>
</tr>
<tr>
<td>Tabatabaei et al.</td>
<td>Case-control study in the Western Australian Bowel Health Study; 567 incident CRC cases and 713 controls</td>
<td>Questionnaires on lifestyle and meat consumption</td>
<td>Amount of red baked meat consumed statistically significant inverse trend of association with CRC (OR 0.73, 95% CI 0.53-1.01). No other statistically significant or meaningful associations with any of the types of meat cooked by any method and CRC risk</td>
<td>Data does not support the hypothesis that meat consumption is a risk factor for CRC</td>
</tr>
</tbody>
</table>
Table 3 Association between nitrates/nitrites found in processed meat and CRC.

<table>
<thead>
<tr>
<th>Author/Topic</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Miller et al. [12]</td>
<td>Case-control study; 989 cases/1,033 healthy controls</td>
<td>FFQ w/a meat-specific module</td>
<td>Significant positive association for nitrates/nitrites and proximal colon cancer (OR 1.57, 95% CI 1.06-2.59)</td>
<td>HCAs, PAHs, nitrates, and nitrates may be involved in CRC etiology</td>
</tr>
<tr>
<td>Parr et al. [15]</td>
<td>Prospective cohort study; population-based Norwegian Women and Cancer cohort; 84,538 women 459 colon and 215 rectal cancer cases with follow-up</td>
<td>FFQ that covered habitual, but not total diet, during the previous year</td>
<td>Processed meat intake ≥60 vs. &lt;15 g/day was associated with significantly increased cancer risk in all subsites.</td>
<td>Did not support an association between CRC risk and intake of red meat or meat cooking methods, but a high processed meat intake was associated with increased risk of proximal colon, distal and rectal cancer</td>
</tr>
<tr>
<td>Dellavalle et al. [17]</td>
<td>Prospective cohort study; cohort from the Shanghai Women's Health Study of 73,118 women age 40-70; 619 CRC cases</td>
<td>77-item FFQ</td>
<td>Nitrate intake was not associated with CRC risk (HR 1.08 95% CI 0.73-1.59). Those with vitamin C intake below the median (83.9 mg/day) and as a result had higher potential exposure to NOCs, CRC risk increased with increasing quintiles of nitrate intake (highest vs. lowest quintile HR 2.45 95% CI 1.15-5.18).</td>
<td>An association between dietary nitrate and nitrate was not observed; however, findings suggest that a high intake of nitrate among subgroups expected to have high endogenous N-nitroso compounds (NOC) exposure (those with low vitamin C intake) are associated with increased CRC risk</td>
</tr>
<tr>
<td>Hu et al. [18]</td>
<td>Case-control study, 19,732 cases</td>
<td>69-item FFQ</td>
<td>Significant results were found for processed meat and CRC (OR 1.5, 95% CI 1.2-1.8)</td>
<td>Findings add to the evidence that high consumption of salt and processed meat may play a role in the etiology of several cancers–stomach, pancreas, lung, prostate, testis, kidney, and bladder</td>
</tr>
<tr>
<td>De Stefani et al. [19]</td>
<td>Case-control study in Uruguay; 6,060 participants (3,528 cases and 2,532 controls)</td>
<td>64-item FFQ</td>
<td>Positive association between processed meat and CRC (OR 2.39, 95% CI 1.76-3.24)</td>
<td>Processed meat intake could be a powerful mutlorgan carcinogen affecting CRC and also the stomach, esophagus, and lungs</td>
</tr>
</tbody>
</table>

These compounds can form during the process of meat curing or within the body from dietary consumption of nitrate/nitrite. Several NOCs are known carcinogens in humans [1]. Based on the convincing evidence of the carcinogenic effect of these compounds, it is recommended to avoid processed meat.

Heme iron in meat

Iron is a potentially toxic, tightly regulated mineral that plays a central role in oxidative metabolism and is a component of several enzymes. A catalytic reaction of free iron can generate free radicals that may lead to oxidative damage of DNA, protein, and membrane lipids of cells; therefore, the body strictly regulates iron transport and metabolism to reduce the likelihood of oxidative damage [20]. Heme iron is found in foods of animal origin, while non-heme iron is found in plant foods. High intakes of red meat may result in more heme iron absorption leading to greater oxidative stress and increasing the potential for DNA damage [1]. In addition, dietary heme can induce colonic cytotoxicity and hyperpolorlferation leading to inflammation [21].
Table 4 Association between heme iron in meat and CRC.

<table>
<thead>
<tr>
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</tr>
</thead>
<tbody>
<tr>
<td>Kato et al. [22]</td>
<td>Case-control study with 1205 cases and 1547 controls between the ages of 45 and 80</td>
<td>Phone interview with structured questionnaires regarding usual diet</td>
<td>CRC was increased by heme iron intake (OR 1.47, 95% CI 1.14-1.89)</td>
<td>Heme iron from meat may increase the risk of CRC</td>
</tr>
<tr>
<td>Gilseng et al. [23]</td>
<td>Prospective cohort study from the Netherlands Cohort Study; 4026 subcohort members, 435 colon and 140 rectal cancer patients</td>
<td>150-item semiquantitative FFQ</td>
<td>Heme iron associated with increased risk of CRC harboring activating gene mutations in KRAS (Kirsten ras) (HR 1.71 95% CI 1.15-2.57) and CRC without truncating mutations in APC (adenomatous polyposis coli) (HR 1.79 95% CI 1.23-2.60)</td>
<td>Positive association found between dietary heme iron and CRC risk, with the suggestion of alkylating rather than oxidative DNA-damaging mechanisms are involved in carcinogenesis of CRC</td>
</tr>
<tr>
<td>Hara et al. [24]</td>
<td>Prospective cohort study in Japan; 39,721 men and 45,376 women age 45-74</td>
<td>138-item FFQ</td>
<td>No association was found linking zinc and heme iron intake with CRC in either Japanese men (HR 1.06, 95% CI 0.79-1.42) or women (HR 0.88, 95% CI 0.61-1.29)</td>
<td>Results in a Japanese population with lower intakes and different major food sources of zinc and heme iron in comparison to the Western diet suggest zinc and heme iron intake not associated with CRC</td>
</tr>
<tr>
<td>Zhang et al. [25]</td>
<td>Prospective cohort study in Nurses’ Health Study and Health Professionals Follow-Up Study; 2,114 incident CRC cases</td>
<td>FFQ</td>
<td>Total iron intake, intake of dietary iron, and supplement uses did not have a significant association with CRC (RR 1.10, 95% CI 0.93-1.30)</td>
<td>Does not support strong roles of zinc and heme iron intake in CRC risk</td>
</tr>
<tr>
<td>Ashmore et al. [26]</td>
<td>Case-control study in Pennsylvania; 1005 incident cases and 1062 controls</td>
<td>Modified FFQ that included supplement use and a meat-specific module</td>
<td>No significant associations between heme iron or total iron intake and CRC incidence (OR 0.70, 95% CI 0.44-1.11) Supplemental iron intake of more than 18 mg/day versus non was positively associated with CRC incidence (OR 2.31, 95% CI 1.48-3.59)</td>
<td>Consumption of more than 18 mg/day of supplemental iron may increase risk for CRC, but heme or total iron was not found to have an association with CRC</td>
</tr>
</tbody>
</table>

Table 4 summarizes five studies conducted after the SER that explore the association of iron with CRC risk. Two studies supported the association, while three of the studies concluded that the evidence is insufficient to confirm the association. Kato et al. [22] conducted a population-based case-control study investigating the associations of dietary iron, smoking, and intestinal bacteria with the risk of CRC. Results showed that CRC was increased by heme iron intake (OR 1.47, 95% CI 1.14-1.89) [22]. The study discussed the importance of heme iron derived from animal meat rather than total iron intake as a risk for CRC. In a 2013 prospective cohort study, researchers found a significant association between CRC risk and heme iron intake activating gene mutations in KRAS2 (a step that may lead to cancer development) (HR 1.71, 95% CI 1.15-2.57) and risk of CRC without truncating mutations in APC (adenomatous polyposis coli) (HR 1.79, 95% CI 1.23-2.60) [23]. Results indicate that an alkylating mechanism may underlie the positive association between dietary heme and CRC risk, versus a DNA-damaging mechanism [23].

Two prospective cohort studies found no association of heme iron intake and CRC risk. A large population-based prospective study examined the association between intake of zinc and heme iron and risk of CRC in a general Japanese population [24]. No association was found linking zinc and heme iron intake with CRC in either Japanese men (HR 1.06, 95% CI 0.79-1.42) or women (HR 0.88, 95% CI 0.61-1.29) [24]. Another study examined zinc and heme iron intake and CRC risk using data from the Nurses’ Health Study and Health Professionals Follow-Up Study, which included 2114 incident CRC cases [25]. Results showed total iron intake, intake of dietary iron, and supplement use did not have a significant association with CRC (RR 1.10, 95% CI 0.93-1.30) [24]. Consumption of more than 18 mg/day of supplemental iron may increase risk for CRC, but heme or total iron was not found to have an association with CRC [25].

KRAS: Also referred to as GTPase Kras and V-Ki-ras2 Kirsten rat sarcoma viral oncogene homolog. This is a protein encoded by the KRAS gene in humans. A mutation may lead to the development of many cancers.

APC: Adenomatous polyposis coli, also known as deleted in polyposis 2.5 (DP2.5). This is a protein encoded by the APC gene in humans. Colorectal cancer may result due to a mutation of this gene CI 1.23-2.60) [23]. Results indicate that an alkylating mechanism may underlie the positive association between dietary heme and CRC risk, versus a DNA-damaging mechanism [23].
incidence of CRC (OR 0.70, 95% CI 0.44-1.11) [26]. Intake of supplemental iron of more than 18 mg/day versus 0 mg/day had a positive association with incidence of CRC (OR 2.31, 95% CI 1.48-3.59), suggesting that consumption of >18 mg/day of supplemental iron may increase CRC risk, although these data do not support the association of total or heme iron intake with risk of CRC [26].

Discussion

This paper summarized new study findings regarding the proposed relationships between red and processed meat consumption and CRC that have been published since the 2011 CUP Report. This topic is particularly relevant given the consumption of red and processed meat in the US and the incidence of CRC. Overall, findings of the updated literature continue to support the WCRF/AICR guidelines and the importance of limiting the consumption of red meat to 18 ounces per week and to minimize processed meat consumption altogether for the prevention of CRC. A dose-response relationship between red and processed meat and CRC lends further credence to these findings reported [27,28], with additional support highlighting the proposed underlying mechanistic roles of HCAs/PAHs, NOCs, and heme iron. To date, the exact role of these compounds in CRC development remains elusive, yet they are clearly important in this diet-disease relationship and warrant further investigation [29].

While the data linking meat and CRC is relatively strong, some inconsistencies are evident between studies and may be attributed to the inherent difficulties of dietary assessment. In the studies reviewed, each employed either a food frequency questionnaire (FFQ) or another form of self-administered questionnaire to collect dietary assessment data (e.g., food record, food diaries). FFQs are a valid and reliable method to measure dietary intake and to assess its relationship with outcomes. They are advantageous and applied widely because they are representative of usual intake, can be used in very large studies, rank participants according to intake for analysis, and provide a cost-effective way to collect critical dietary data [30]. Despite these advantages, FFQs are prone to error due to recall bias and the subjective nature of the assessment tool [30]. Recall bias is an inherent limitation, as FFQs rely on the participants’ memory over a specified period of time, typically several months to a year. Some degree of measurement error can be attributed to the subjective nature of these tools, since people have the tendency to underreport what they have eaten, especially when it requires recording. In addition, participants may inaccurately account what they have eaten in the past with response distortion of what they consider healthy versus unhealthy [31]. Because FFQs depict usual intake they are considered less accurate when assessing absolute intake of a specific food group or nutrient, such as meat consumption in these studies. Further, several studies used different FFQs or other comparable dietary assessment questionnaires. While most appropriate for the population under study, these methodological differences pose limitations for the direct comparison of responses and outcomes assessment across studies. All of these previous investigation utilized statistical methods to control for potential confounders, including age, sex, smoking, alcohol intake, body weight, BMI, and family history; however, residual confounding may still bias or impede our abilities to depict more precise diet-disease relationships due to the natural variations in dietary intake not detected by these dietary assessment methodologies. Finally, FFQs and other self-administered dietary assessment methods do not take into consideration several influencing factors, such as cooking methods or temperature, which are now recognized as important considerations in CRC development [32].

While the negative associations between red and processed meat consumption and CRC continue to build, a broad body of evidence supports that plant-based dietary patterns, fruit and vegetable consumption, and specific plant constituents may reduce the overall risk of cancer, as well as cancer recurrence. Several studies have identified dietary patterns to investigate the association of the total diet on CRC risk. While the study designs, methods, participant populations, and derived dietary patterns differ, the three reviews [33-35], three case-control [36-37], and two prospective cohort studies [38-40] all concluded that meat-based diets are associated with an increased risk of CRC and that plant-based diets, rich in fruit, vegetables, and low in red and processed meat, may reduce CRC risk. These findings are consistent with Tantamango-Bartley et al. [41] who found that vegetarian versus non-vegetarian dietary patterns were associated with significantly reduced cancer incidence (HR 0.92, 95% CI 0.85-0.99). This study also reported that vegan dietary practices were even more protective for overall cancer incidence (HR 0.66, 95% CI 0.47-0.92) [41]. Lanou and Svenson [42] conducted an analysis of recent reports on cancer risk in vegetarians and found that the direct and indirect evidence supported that vegetarian diets are useful in reducing cancer risk. Findings from a systematic review and meta-analysis exploring cancer incidence in vegetarians supported that vegetarians have significantly lower overall cancer incidence than non-vegetarians (RR 0.82, 95% CI 0.67-0.97) [43]. Although there is a cause and effect relationship between red and/or processed meat consumption and CRC, strong evidence exists to support a plant-based diet approach for overall cancer prevention.

In support of the WCRF/AICR recommendations, many professional organizations also promote a plant-based diet and limiting red and processed meat consumption for the general prevention of cancer, not specifically CRC. The Academy of Nutrition and Dietetics (AND) position paper on vegetarian diets reports that appropriately planned vegetarian diets are nutritionally adequate and are associated with a lower body mass index and a reduced incidence of cancer overall [44]. The American Cancer Society (ACS) supports that the most important ways to reduce cancer risk include maintenance of a healthy weight, regular physical activity, and consuming a mostly plant-based diet limiting saturated fat [45]. The ASC guidelines specifically recommend limiting red and processed meat, while consuming at least two and a half cups of fruit and vegetables each day [45]. The World Health Organization (WHO) also states that dietary habits including diets high in fruits and vegetables may have a protective effect against a multitude of cancers,
while consuming excess red and processed meat may have an association with increased CRC risk [46,47]. Although there are several different recommendations by these authoritative groups, it is clear that dietary patterns (e.g., plant based) and dietary intake (e.g., limited meat consumption) are the cornerstone of cancer prevention.

Conclusions

With high cancer incidence rates and mounting evidence regarding the role of diet in cancer prevention, dietary intervention in now more than ever crucial to help reduce the number of new cancer cases each year. The 2011 CUP Report supports the AICR guidelines that recommend limiting red and processed meat and consuming a plant-based diet for primary cancer prevention. Evidence since the report continues to reaffirm the relationship between red and processed meat and CRC risk. The proposed mechanisms underlying the association include HCAs/PAHs, N-nitroso compounds, and heme iron; however, these relationships are far from definitive. All of these compounds are implicated in the carcinogenesis process and the strongest association to date is found for CRC. A plant-based diet not only minimizes compounds hypothesized to increase cancer risk, but emphasizes high intakes of fruits and vegetables to increase to increase the consumption of protective compounds found in plant-based foods. Based on the continually emerging evidence, the ideal recommendation for cancer prevention involves making healthy lifestyle choices. Included in this recommendation is maintaining a healthy weight, engaging in regular physical activity and consuming a diet that emphasizes plant sources, reduced consumption of red and processed meat, and increasing plant-based whole foods.
References


46 http://www.who.int/ena/titles/fruit_vegetables_ncds/en/