Young Adult Cancer: Influence of the Obesity Pandemic

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Objective: The purpose of this article is to review the association of the obesity pandemic with appearance of cancers in young adults under age 50 and to define potential mechanisms by which obesity may accelerate the development of malignancy.

Methods: A comprehensive narrative review was performed to integrate preclinical, clinical, and epidemiologic evidence describing the association of obesity with cancer in young adults based on a search of PubMed and Google databases.

Results: Results from more than 100 publications are summarized. Although they differ in age groups analyzed and incidence of obesity, sufficient data exists to suggest an influence of the obesity pandemic on the increase of cancer among young adults.

Conclusions: Cancer in young adults is occurring with increasing frequency. Overweight and obesity have become major public health issues reaching pandemic proportions. Excess weight is associated with increased cancer risk, morbidity, and mortality. Multiple murine models indicate that obesity not only increases cancer incidence but also accelerates its development. Thus, the possibility exists that overweight and obesity may be contributing to the appearance of specific malignancies at younger ages. This prospect, in association with the worldwide expansion of obesity, suggests an impending explosive increase in obesity-associated cancers in young adults.

Introduction

Cancer in young adults is being reported with increasing frequency and has become a matter of urgent concern (1). At the same time, overweight and obesity have become major public health issues in both children and adults, reaching pandemic proportions worldwide (2,3). While it has been clearly documented that excess weight is associated with both increased risk of occurrence and increased morbidity and mortality for multiple malignancies (4-6), there has been relatively little focus on the impact of overweight and obesity on shifts in timing of cancer appearance to individuals of younger age. However, recent Centers for Disease Control and Prevention data indicate an increase in overweight- and obesity-associated cancers in 20- to 49-year-old individuals (Supporting Information Supplement S1). Importantly, multiple murine models indicate that obesity and obesogenic diets not only increase the incidence of malignancy but also accelerate its development and shift its occurrence to earlier ages (5,7-17).

Thus, the possibility needs to be considered that overweight and obesity may be contributing significantly to the clinical appearance of some malignancies at younger ages. This prospect, in association with the continued worldwide expansion of obesity (2,3), suggests an impending explosive increase in obesity-associated cancers in young adults. Anticipation of the potential dire consequences of this evolution compels careful epidemiologic monitoring; more research on mechanisms by which obesity promotes and accelerates cancer, especially in young adults; development of focused strategies for prevention; and potentially new approaches to screening and care.

The goals of this article are 1) to enhance awareness of the obesity–cancer linkage; 2) to illustrate how both obesity and obesogenic diets may shift appearance of obesity-promoted cancers to younger age groups, especially into the 20- to 50-year-old age group; 3) to examine preclinical murine models and potential mechanisms by which obesity and obesogenic diets accelerate the appearance of malignancy; 4) to review the epidemiologic and clinical evidence indicating where this may already be happening; 5) to identify which obesity-associated cancers are most likely to pose this threat; and 6) to consider approaches to better document and avert the crisis.
Obesity–Cancer Linkage

Although adolescent and young adult cancers have been operationally defined as those occurring in the 15- to 39-year-old age group (18), this article is focused on malignancies, most commonly associated with patients over age 50, that have recently been reported with increasing frequency in the younger-than-50-year-old age group. Moreover, because this article examines the impact of obesity on cancers in young adults, it will concentrate on the 13 cancers listed in Table 1, which, based on epidemiologic review by the International Agency for Research on Cancer (IARC), have been identified as having sufficient evidence to define an association with obesity has not reached the level of significance as those recently reported (6). This article uses the following categories of BMI (weight in kilograms divided by square of height in meters): normal weight, BMI = 18.5-24.9; overweight, BMI = 25.0-29.9; obesity, BMI ≥ 30; and severe or morbid obesity, BMI ≥ 40 (6).

Table 1, column 1 lists the 13 tumors recently reported by the IARC for which there is sufficient evidence to define an association with obesity and specific malignancies (6). These malignancies are arranged in order of annual incidence of new US cases (column 2) (19,20). Column 3 lists the US population attributable fraction (PAF) as the percentage of each malignancy attributed to obesity for both males and females (21). These data demonstrate an important contribution of obesity to cancers of colon and rectum, thyroid, esophagus, pancreas, and kidney in men and to breast, colon, kidney, endometrium, esophagus, and gallbladder cancers in women. With 253,000 new cases of breast cancer in the United States, a 14% PAF calculates to 35,420 new cases per year attributable to obesity. For colorectal cancer (CRC), adjusting for male/female distribution, the PAF indicates 22,655 new cases of CRC in men and 10,812 new cases in women attributable to obesity. Applying similar calculations to the incidence and PAF data provided in Table 1 indicates that in 2017, more than 144,000 of those cancers occurring in the United States were attributable to obesity. However, this number is probably an underestimate, as PAFs are not available for several obesity-associated malignancies such as liver, myeloma, gastric cancer, or meningioma.

The fourth and fifth columns indicate the peak and usual age range incidence for each of these tumors. The median age at which all cancers are diagnosed in the United States is 66 years. While most are diagnosed in patients older than 50 years (20), the appearance of thyroid, ovarian, endometrial, and CRC cancers and meningiomas is not uncommon in patients younger than 50 (20). Strikingly, as shown in the sixth column of Table 1, of the 13 IARC obesity-associated malignancies, at least 9 (shown in bold) have been reported as occurring in young adults and are in the top 20 adolescent and young adult cancers (18). The last column indicates murine

### Table 1: Relation of obesity-associated cancers to young adult malignancies and murine models

| Obesity-associated cancer | US incidence × 10^-3 | Population attributable fraction %, M/F (21) | Peak age incidence, years | Usual age range of all years with incidence >15% | Percent new cases in 20-44 years age group | DIO and HFD in murine models promoted cancer |
|---------------------------|----------------------|---------------------------------------------|--------------------------|------------------------------------------|----------------------------------------|-----------------------------------------------|
| Breast                    | 253                  | -/14                                       | 62                      | 55-84                                    | 10.5                                   | MMTV-TGFbeta (7)                             |
| Colon and rectal          | 135                  | 32/17                                      | 67                      | 45-84                                    | 5.8                                    | APCKn (8)                                     |
| Kidney                    | 63                   | 25/34                                      | 64                      | 55-74                                    | 7.8                                    |                                               |
| Endometrial               | 61.3                 | -/48                                       | 62                      | 45-74                                    | 7.3                                    | Pten+/−/16 (11)                               |
| Thyroid                   | 57                   | 32/5                                       | 51                      | 20-64                                    | 23.9                                   | Thrb^V/V, Pten+/−/16 (13)                     |
| Pancreas                  | 54                   | 14/11                                      | 70                      | 55-84                                    | 2.4                                    | Kras^G12D, conditional (9,10)                |
| Liver                     | 41                   | NA                                         | 63                      | 55-84                                    | 2.5                                    | C57BL/6j (11) MUP-UPA (12)                   |
| Myeloma                   | 30                   | NA                                         | 69                      | 55-84                                    | 3.5                                    | KwLwRij (14)                                 |
| Gastric cardia            | 28                   | NA                                         | 68                      | 55-84                                    | 6.2                                    |                                               |
| Meningioma                | 27                   | NA                                         | 58                      | 45-74                                    | 16.8                                   |                                               |
| Ovary                     | 22                   | -/7                                        | 63                      | 45-84                                    | 10.6                                   | Kp8 (15)                                     |
| Esophageal adenocarcinoma | 17                   | 44/48                                      | 67                      | 55-84                                    | 2.3                                    | L2-IL-1beta (17)                             |
| Gallbladder               | 7                    | -/53                                       | 85±9                    | 65-90                                    |                                               |                                               |

Notes:
- Obesity-associated cancers identified by 2016 IARC analysis (6).
- US incidence of specific cancers from American Cancer Society Cancer Facts & Figures, 2017 (19).
- Peak age incidence from SEER Cancer Statistics, 1975-2014 (20).
- Usual age range years from SEER Cancer Statistics (20) combining all decades with incidence ≥15% for each malignancy.
- Percent new US cases in 20- to 44-year-old age group from SEER Cancer Statistics (20) combining 20- to 34- and 35- to 44-year-old, age group. Values in bold font indicate malignancies among top 20 invasive cancers in the United States at ages 20 to 39 years (18).
- Age range for meningioma provided for all primary brain tumors (19).
- Gallbladder age incidence and range from UK data, 2015 (22).
- NA, not available.
model systems in which nine of the malignancies have been shown to be accelerated and become more aggressive in association with obesity (7-17).

**Obesity Accelerates Cancer Development**

From a mechanistic viewpoint, overweight and obesity are generally considered to be promoters of cancer progression (5). Thus, overweight and obesity promote cancer by multiple concurrent mechanisms, including 1) stimulation of low-grade inflammation and oxidative stress with increased levels of proinflammatory cytokines such as IL-6, tumor necrosis factor (TNF), and increased reactive oxygen species (ROS), the latter of which may also contribute to mutagenesis; 2) alteration of growth-promoting factor levels, especially insulin and insulinlike growth factor (IGF-1), which increases in association with metabolic syndrome and insulin resistance; 3) altered sex steroid hormones with increased conversion of androgens to estrogens resulting from increased adipose tissue production of aromatase, the enzyme responsible for this conversion; 4) altered adipocytokine proteins, including increased growth-promoting and pro-inflammatory components such as leptin, retinol binding protein 4, resistin, and visfatin and reduced growth-controlling adipokines such as adiponectin; 5) alterations in intestinal microbiome with expansion of tumor-promoting species such as fusobacteria; and 6) mechanical effects of obesity such as those leading to hiatal hernia and gastroesophageal reflux disease, predisposing to esophageal adenocarcinoma (5).

Figure 1 provides a conceptual model, supported by multiple murine studies, of how obesity impacts cancer by accelerating its development (7-17). As postulated for development of colon cancer, and now widely accepted for multiple malignancies, mutations in a gatekeeper gene (13,22,23), sometimes similar to those mutations causing hereditary cancer syndromes, initiate sporadic tumors. Mutated cells then progress through a multistage process in which multiple genetic changes ultimately lead to the development of a benign premalignant, and then to malignant, neoplasm with invasive and subsequently
metastatic properties. For CRC, transformation from normal epithelium to benign adenoma, ultimately leading to frank cancer and metastatic disease, is projected to require at least seven independent genetic events and possibly mutation in as many as 15 driver genes (24) as well as multiple epigenetic alterations (25). In some cases, this process may require long latent periods extending to multiple decades to progress from normal epithelium to frank cancer.

The rate of progression to invasive cancer is determined by multiple factors, including mutation and proliferation rates, which are affected by DNA damage response, DNA repair systems, and a host of growth factors. This rate may vary among different tumors and even among different transformed clones in the same individual. Thus, obesity may enhance mutation rates by generation of increased ROS. Importantly, however, high-fat diets (HFDs) and diet-induced obesity (DIO) have been shown to accelerate tumor growth rates in association with production of increased growth factors, such as insulin, IGF-1, leptin, retinol binding protein 4, and others (8,26,27).

As shown in Figure 1, development of obesity, usually due to a combination of HFD and decreased physical activity, results in expanded fat mass, characterized by increased number and size of adipocytes, some of which undergo necrosis and become surrounded by macrophages to form crown-like structures with a propensity for releasing proinflammatory cytokines such as IL-6 and TNFα (28). In addition, expanded and inflamed adipose tissues may provide increased levels of multiple growth-promoting cytokines, adipokines, and hormones, many of which accelerate the multistage transition from normal tissue to invasive and metastatic cancer. This process not only accounts for the accelerated development of tumors in the presence of adipose tissue excess, but it also explains why patients with obesity-driven cancers may present with more advanced tumors at earlier ages. Thus, the long latent period required for initial presentation of many tumors provides the basis for obesity to impact the process and, in fact, accelerate both the appearance and extent of clinical disease. Accordingly, it is expected that initial mutations for sporadic cancer will occur with similar frequency and at similar ages in both normal patients and those with obesity. In addition, it is possible that obesity-associated inflammation and ROS may further contribute to mutagenesis and cancer initiation. Nonetheless, the metabolic and growth-promoting consequences of concurrent or prior obesity can provide the stimulus for accelerated development of cancer and associated comorbidities, including death.

**Clinical and Epidemiologic Evidence Indicating That Obesity Shifts Malignancies to Younger Ages**

From a clinical and epidemiologic viewpoint, we focus initially on CRC, which has become one of the major adult tumors generating alarm for its increasing appearance in young adults (33). CRC, usually occurring between 45 and 84 years of age with peak incidence at 67 years (20), and uncommonly seen in young adults, is now being increasingly identified in both men and women below age 50 (20,34-39), with greater increase noted for left-sided sigmoid and rectosigmoid CRC than right-sided CRC (35).

Analysis of Surveillance, Epidemiology, and End Results Program (SEER) Population data and multiple Hospital Based Cancer Registries, covering periods from 1973 to 2017, indicates that CRC incidence has remained stable and/or decreased in people over 50 by as much as 3% per year. In contrast, CRC has shown an average 1.5% increase per year among 20- to 40-year-old men and women (40,41). Moreover, younger patients have been noted to present with more advanced, higher-stage, more poorly differentiated disease, and those presenting with stage IV CRC have shown inferior survival.

The overall decrease in incidence of CRC has been attributed to expanded screening programs and removal of early premalignant adenomas. Because of the much higher incidence of CRC in older individuals, these programs have been primarily targeted at patients over 50. Thus, increase in incidence and more advanced stage at presentation among young adults have been attributed, in part, to lack of screening and to tumor promotion by lifestyle factors including obesity, consumption of red and processed meat, and possibly alcohol and tobacco use (35,42).

Many of the above reports point to a concurrent increase of obesity and CRC in the young. Some have documented increased obesity and cancer in the same population; however, few have provided systems demonstrating that obesity promotes and accelerates development of malignancies

The shortening of the latent period from benign to malignant disease in association with obesity has been most clearly demonstrated at the clinical level, where disease-associated monoclonal immunoglobulin provides a biomarker for early detection and demonstration that obesity accelerates the conversion of monoclonal gammapathy of unknown significance (MGUS) to multiple myeloma (MM) (29). In further support of this proposal that obesity does not initiate but rather promotes cancer progression, almost all murine models in which obesogenic diets and DIO promote tumors require experimental utilization of genetically modified animals containing cancer-predisposing genes or transplantation of preexisting tumor cell lines (5,7-17). It is noteworthy that in some murine models, HFD has been shown to promote CRC and breast cancer in mice that are resistant to DIO (7,8,30). These studies indicate that proinflammatory and growth-promoting effects of HFD, even in the absence of DIO, may accelerate tumor progression. Other murine systems have shown that even after HFD-induced DIO and subsequent weight loss, the tumor-promoting effects of obesity may endure for varying time periods, thereby providing a model for promotion of adult tumors by childhood, adolescent, and young adult obesity (7,8,30).

The contribution of proinflammatory and growth-promoting factors as mediators in the HFD- and DIO-accelerated malignancies is further illustrated by the demonstration that tumor-promoting effects of obesity can be abrogated by molecular or pharmacologic interference with proinflammatory and growth-promoting pathways such as pharmacologic inhibition of receptors for insulin or IGF-1 (31), molecular interference with leptin receptor (32), and genetic and pharmacologic interference with proinflammatory activity of complement system (8). It is further noteworthy that not all HFDs are equal in promoting malignancy, as shown by olive oil, an important component of the Mediterranean diet, which was found to protect against HFD acceleration of gastrointestinal neoplasia in APCMin mice (8).
obesity demographics in the CRC patients (38). In addition, although not specific for young adults, some series indicate an association of obesity with increased risk of sigmoid and rectosigmoid cancers (43). In an important study of more than 1.1 million Israeli Jewish men with 19.5 million person-years of follow up, overweight and obesity in adolescents aged 16 to 19 years were associated with a substantially increased risk for colon cancer (HR = 1.53, 95% CI: 1.17-2.0) but not for rectal cancer in adult years. The median age for patients with newly diagnosed colon cancer was 43.3 ± 8.7 years, thereby supporting an association of adolescent-detected obesity with young adult colon cancer (37) and suggesting effects of obesity over a long latent period of cancer development.

In addition to its association with increased risk for CRC, obesity is also associated with a twofold increase in risk for colorectal adenoma (CRA), a premalignant precursor to CRC (44). CRAs have been commonly reported in patients younger than age 55 (45) and have been noted to be more advanced in patients with obesity (46). In a study of people examined across an age range from 30 to greater than 70 years, high BMI was identified as a risk factor for CRA in 30- to 39-year-old men and 40- to 49-year-old women (36). Thus, subjects with overweight or obesity are at increased risk for developing CRC and its precursor, CRA, during young adulthood. Moreover, obesity has been shown to precede the diagnosis of CRA and CRC by long latent periods (47).

In summary, the clinical development of CRA and CRC fits well with the model provided in Figure 1, including increased and early development of obesity-associated benign adenoma preceding cancer with a long latent period, providing time for the impact of obesity-stimulated growth factors to accelerate tumor development. Moreover, in addition to decreased screening in young adults, obesity-promoted progression may help explain why CRC in the young is more advanced at the time of presentation (33).

While this discussion has focused on sporadic colorectal neoplasia and its increasing appearance in young adults, additional insight is provided by patients with known inherited predispositions to CRC, including familial adenomatous polyposis or hereditary non-polyposis colon cancer (HNPPC), bearing, respectively, mutations in the gatekeeper genes, adenomatous polyposis coli, or mismatch repair genes (MLH1, MSH2, MSH6, PMS2, EPCAM) (48). These patients commonly develop CRA or CRC at younger ages, when CRC risk has been shown to be increased in association with obesity (49,50). In a study of 937 HNPPC carriers followed at 14 institutions, with median age enrollment 44.9 years (36-53 years), obesity was associated with a 2.4-fold greater risk for CRC compared with normal and underweight reference groups. Interestingly, there was no increase in risk in HNPPC patients with obesity randomly assigned to aspirin (ASA), 600 mg daily, suggesting that obesity-promoted CRC in HNPPC patients may be reduced by regular ASA use (51).

Female breast cancer is the most common US malignancy included on the IARC obesity-associated cancer list, with peak incidence at 62 years and usual age range 55 to 84 years (20). Breast cancer in postmenopausal women is usually estrogen receptor positive and is associated with increased risk in association with obesity (4). Of the tumors listed in Table 1, breast cancer is unique in that a major variety, premenopausal breast cancer, characterized by estrogen receptor negative status, has been noted to occur at a relatively constant rate of 40% by age 40 years (52), and obesity is associated with an overall decreased risk of premenopausal breast cancer (53). Thus, because of the already significant occurrence of premenopausal breast cancer in patients under 40 years of age, it is difficult to determine if there is an obesity-associated shift to younger age. However, premenopausal women at high risk for breast cancer, including prior history of lobular carcinoma in situ, generally considered a multifocal premalignant precursor, have shown significantly increased risk of developing breast cancer in association with obesity (54). These high-risk premenopausal women fit well into the latent process depicted in Figure 1 that is accelerated by obesity. Moreover, unique insight is provided by patients with triple negative breast cancer (TNBC), tumors that lack expression of estrogen receptor, progesterone receptor, and human epidermal growth factor receptor 2 (55). These tumors, commonly identified in premenopausal women, are refractory to hormonal and cytotoxic chemotherapy (55). In a retrospective review of invasive breast cancer among 1,064 patients from Walter Reed National Military Medical Center, 160 patients had TNBC, of whom 89 were below age 50 and 148 either had overweight or obesity. Thus, TNBC in all patients, including those younger than age 50, is highly associated with obesity (56).

Further insight is provided by patients with hereditary breast and ovarian cancer wherein 80% of cases have been attributed to mutations in BRCA1/BRCA2 (57). In a series of 176 multigenerational kindreds, earlier appearance of breast cancer was noted in successive generations, with age at diagnosis shifting from 51.8 years in grandparents to 48.7 years in parents, followed by 41.9 years in probands and 34.7 in children (58). The shift to earlier age in successive generations was attributed to lifestyle factors including obesity. In other studies, weight gain and number of pregnancies have been shown to significantly increase risk of breast cancer, whereas weight loss between age 18 and 30 years has been associated with decreased risk of breast cancer (59).

Ductal carcinoma in situ (DCIS), with peak incidence during age 60 to 74 years, is a noninvasive breast cancer precursor and is unusual before age 35. DCIS has recently undergone a marked increase in detection and occurrence from an incidence of 1.8 per 100,000 in 1973-1975 to 32.5 in 2004. The increase has mainly occurred in patients older than 50; however, DCIS has been noted to be increased in all age groups in association with obesity (60). Thus, the breast cancer precursor DCIS, familial breast cancer associated with BRCA mutation, and TNBC are all increasing in incidence in young adults, and the increases are associated with obesity (54-60).

Renal cell carcinoma (RCC), the third most common IARC obesity-associated malignancy (6), has a peak incidence for diagnosis at age 64 (20); however, several retrospective series have reported RCC patients younger than 45 (61). Although up to 60% of patients with RCC have been reported as having overweight/obesity (62), the specific percentage of patients under 45 with excess weight has not been reported. However, in a case control study of 1,214 patients and 1,234 controls, early adult obesity was associated with a 60% increase in risk for RCC (63).

Endometrial cancer in the United States is the most common malignancy of the female genital tract. It is most frequently diagnosed in postmenopausal women aged 45 to 74 years (19,20). However, 2% to 14% of endometrial cancers have been reported to occur in women aged 40 years and younger (64). Sporadic endometrial
Obesity: Young Adult Cancer

Obesity-mediated liver damage progresses through nonalcoholic fatty liver disease (NAFLD) showing steatosis and lipid deposition in liver cells without inflammation, proceeding to nonalcoholic steatohepatitis characterized by further fatty acid deposition and ballooning degeneration of hepatocytes with inflammation, leading to fibrosis, cirrhosis, and HCC (72). In Western countries, where overweight and obesity are common, NAFLD is present in 20% to 40% of the general population (72). The increase in obesity and its comorbidities, including diabetes and metabolic syndrome in Western countries, is projected to have been present for latent periods in excess of 20 years (82). Obesity has been shown to be associated with increased risk for MGUS in women (83). Interestingly, although it is considered a premalignant condition, MGUS shares many of the genetic and cytogenetic changes noted in MM, including activation of c-myc, del(17p), t(4;14), and 1q gains (81). In a retrospective study of 7,878 MGUS patients identified through the US Veterans Health Administration database, 39.8% had overweight and 33% reported BMI. In a pooled analysis of 242 MM cases in patients younger than 50 compared with 1,758 age-matched controls, patients showed a significant positive association of elevated BMI with risk of MM and a greater than twofold increase in MM risk for patients with severe obesity (78). Moreover, the incidence of MM has been noted to be increased in patients who reported heavy compared to lean body shapes during childhood and adolescence (75), providing support for a potentially long latent period for the impact of obesity on malignancy development. Interestingly, chromosomal abnormalities characteristic of MM have been shown to be no different in patients above or below age 45 (79).

Liver cancer commonly occurs in young adults, with 28% of new diagnoses in the 20- to 40-year-old age group. Its incidence in patients under 65 is increasing (65), extending down to the 45- to 49-year-old age group in some South American countries (66). In contrast with many tumors discussed in this article, thyroid cancer in young adults is usually curable because it most frequently is detected early as an asymptomatic neck mass. Cases in young adults show a female preponderance; appear to be differentiated, with papillary histology being more frequent than follicular; and frequently harbor a mutation in the RAS-RAF-MEK-ERK, mitogen activated protein kinase pathway (65).

Pancreatic cancer, with peak incidence in 70- to 80-year-old individuals, is uncommonly observed below age 45 and is increasing in frequency (19,20). Individuals with overweight or obesity between ages 20 and 39 years had 2 to 6 years’ earlier onset of pancreatic cancer compared with normal-weight controls (67). A UK survey conducted between 1998 and 2006 showed no change in incidence rate for males under 50 years; however, a slight increased incidence was reported in females in the 20- to 39-year-old group (68). In a retrospective review covering 1993 to 2008, 33 patients (5.7%) were identified in the 50 years or younger age range. Only 3 (9%) had obesity compared with 4 (12%) of the matched controls (69).

Hepatocellular cancer (HCC), with overall incidence in Western countries peaking at 60 to 70 years of age, is one of the most common cancers on a worldwide basis (70). Incidence rates in the United States have increased by 2.5- to 3-fold over the past 35 years (71). On a global basis, HCC is associated with liver injury from different etiologies including viral infections with hepatitis B and hepatitis C, hepatotoxins including aflatoxin, chronic alcohol abuse, and metabolic alterations that occur with obesity, the latter leading to metabolic syndrome, consisting of obesity, diabetes, insulin resistance, and dyslipidemia (72). The common pathway by which each of these insults leads to HCC includes liver damage, followed by inflammation, usually leading to cirrhosis and then HCC, thereby providing an extended latent period for obesity-promoted carcinogenesis.

Because NAFLD is a predisposing risk factor for HCC, it is noteworthy that NAFLD is increasing in young adults in association with the increased incidence of obesity and metabolic syndrome (74). Thus, NAFLD in young adults aged 18 to 50 years old has increased 2.5-fold over the past three decades and is reported to be present in more than half (57.4%) of young adults with morbid obesity (74).

In a recent US study, the Liver Cancer Pooling Project, composed of 14 separate cohorts and containing 2,087 cases with prospectively measured BMI and waist circumference, showed that excess weight at the time of enrollment was associated with liver cancer in a dose response manner (71). Although most of the cohorts enrolled “older” Americans, 85 patients were younger than 50 years at the time of enrollment, and 18 of these were diagnosed with liver cancer before age 50, 3 of whom had obesity at the time of enrollment (P. Campbell and C. Newton, personal communication). This observation that 18 of 85 patients developed liver cancer before age 50, along with the high incidence of NAFLD in young adults with obesity, indicates the importance of careful surveillance of HCC as another malignancy likely to increase in young adults.

MM, characterized by malignant proliferation of plasma cells, anemia, elevated levels of a circulating monoclonal immunoglobulin, destructive bone lesions, and renal failure, is the second most common hematologic malignancy in the United States and the only primary malignancy of blood cells included by the IARC as related to obesity (6). MM is diagnosed with a peak incidence of approximately 69 years and has maintained a constant incidence for at least the past three decades (75). However, in three series reported since 1992, MM has been reported in patients younger than 45 with incidences of 2.2%, 9.6%, and 15% (76,77). None of these series reported BMI. In a pooled analysis of 242 MM cases in patients younger than 50 compared with 1,758 age-matched controls, patients showed a significant positive association of elevated BMI with risk of MM and a greater than twofold increase in MM risk for patients with severe obesity (78). Moreover, the incidence of MM has been noted to be increased in patients who reported heavy compared to lean body shapes during childhood and adolescence (75), providing support for a potentially long latent period for the impact of obesity on malignancy development. Interestingly, chromosomal abnormalities characteristic of MM have been shown to be no different in patients above or below age 45 (79).

MGUS, characterized by restricted proliferation of a predominant clone of plasma cells, not exceeding 10% of marrow cells and absence of diagnostic criteria for MM (80), is considered a universal premalignant precursor of MM with variable rates of progression (81). MGUS, identified in large population screenings by detection of circulating monoclonal immunoglobulins, is most common in the 80- to 96-year-old age group; however, it has significantly been reported in patients younger than age 50. In some of these cases, it has been projected to have been present for latent periods in excess of 20 years (82). Obesity has been shown to be associated with increased risk for MGUS in women (83). Interestingly, although it is considered a premalignant condition, MGUS shares many of the genetic and cytogenetic changes noted in MM, including activation of c-myc, del(17p), t(4;14), and 1q gains (81). In a retrospective study of 7,878 MGUS patients identified through the US Veterans Health Administration database, 39.8% had overweight and 33% had obesity. Moreover, risk of transformation of MGUS to MM was increased with obesity and black race (29,83).

Esophageal adenocarcinoma (EAC) and gastric cardia adenocarcinoma, both malignancies of glandular epithelium originating near the gastroesophageal junction, have undergone a rapid increase in
incidence over the past two to three decades (84,85). They are consistently associated with overweight/obesity, and 10% of patients presenting with EAC are noted to have morbid obesity (86). Although EAC has a peak incidence in the 80-year-old age group (87), in a retrospective study of 374 patients treated for EAC between 2000 and 2007, 63 (16.8%) were under age 50 (86). EAC may be preceded by a premalignant precursor, Barrett’s esophagus (BE), commonly seen in younger patients, where it is associated with chronic inflammation, gastroesophageal reflux disease, and obesity (87).

In addition to their association with obesity at time of diagnosis, occurrence of both EAC and gastric cardia adenocarcinoma in later years is increased following elevated BMI during early adulthood (age 20) and with progressive weight gain between ages 30 and 50 years (73). In addition to the overall increase in occurrence of BE in young adults (87) there has been a notable increase in EAC in patients under age 40, and more than 10% of patients undergoing surgery for EAC are reported to be ≤50 years old (88).

Meningioma constitutes 20% to 30% of all intracranial neoplasms, with peak incidence in men in the 60- to 69-year-old age group and in the 70- to 79-year-old age group in women (89,90). Cranial irradiation and obesity are risk factors (89), sometimes with long latent periods of more than 20 years between radiation and diagnosis of meningioma. Meningioma in the pediatric age group is seen as part of hereditary syndromes such as neurofibromatosis with inherited NF2 mutations. Somatic mutations of the NF2 gene are frequently identified in sporadic cases of meningioma (90). In a report of 35 patients from a single institution and meta-analysis of more than 450 patients, meningioma occurring during first three decades of life, with an average age at diagnosis of 25 years, had a female predominance but no notation of occurrence of obesity (90).

In contrast to many of the other cancers discussed in this article, epithelial ovarian cancers are not uncommon among young women, in whom they are thought to coincide with activity of the female reproductive cycle (19,20). Increased BMI is a risk factor for epithelial ovarian cancers, and elevated levels of IGF-1, which frequently accompany obesity, are thought to contribute (91). Mutation of mismatch repair genes including germ line mutations occur in a small percentage of patients under 40.

Gallbladder cancer, a rare malignancy in the United States with peak incidence in the 80-year-old group, is rarely seen or reported in patients under 50 years of age (92). Risk factors for gallbladder cancer include obesity and chronic inflammation associated with gallstone disease. Chronic inflammation of 15 or more years has been estimated to result in gallbladder cancer in genetically predisposed individuals. Treatment of cholelithiasis and cholecystitis by surgical removal of the gallbladder has significantly decreased the incidence of gallbladder cancer (93).

Association Between Obesity-Linked and Young Adult Cancers

Of the 13 cancers identified by the IARC as being associated with increased body fat (6), most have their highest incidence rates in older adults. However, 5 of the 13 obesity-associated cancers, including breast, thyroid, uterus, ovary, and stomach cancer, have been identified by US SEER data as occurring in the top 20 cancers in 20- to 39-year-old females and 5 of the 13, including colorectal, thyroid, kidney, stomach and liver cancer, have been identified in the top 20 in 20- to 39-year-old males (18). Of the 13 IARC obesity-associated malignancies, all but gallbladder cancer have been well documented to occur in significant numbers in patients under 50 years of age, and 4 of these malignancies—colorectal, breast, thyroid, and possibly pancreatic cancer—have been reported to be increasing in the female young adult population. Moreover, five premalignant precursors, including CRA for CRC, BE for EAC, NAFLD for HCC, DCIS for breast cancer, and MGUS for MM, have been reported to be increasing in the young adult population in association with obesity. In addition, excess body weight and/or weight gain has been noted to precede presentation of these malignancies by long latent periods, in some cases by multiple decades.

In summary, many of the malignancies noted to occur with increasing frequency in young adults are among the 13 obesity-associated cancers. With the expanding worldwide incidence of overweight and obesity in children and young adults, the long latency period associated with many sporadic cancers, the demonstration in humans and animal models that obesity accelerates the development of cancer, and the probability that even obesity at young age has a long-term effect on tumor progression, it is highly likely if not imminent that obesity will lower the age of occurrence across the age spectrum, shifting multiple malignancies to younger age groups in general and to the young adult population in particular.

Overall, this assessment is limited because many of the reports of malignancies in young adults do not provide anthropomorphic measurements. Moreover, evaluation of these data often underestimates obesity because patients with advanced malignancy frequently present for evaluation after significant weight loss. Further evidence of an association of cancer and obesity in young adults will require more consistent reporting of anthropomorphic data at time of diagnosis as well as premorbid data when available. Because body mass may be considerably reduced at time of cancer diagnosis, it is important that these data be monitored in a prospective fashion among the healthy pediatric and young adult population and made available for analysis if and when malignancy is diagnosed. However, documenting and reporting this information are critically important to more firmly establish the relation of obesity to young adult cancers.

Disrupting the Linkage Between Obesity and Young Adult Cancers

The data cited in this article portend an imminent threat of the impact of the obesity pandemic on an age shift in occurrence of obesity-associated malignancies, including their appearance in young adults. This occurrence will require increased cooperation between adult, adolescent, and pediatric oncologists, endocrinologists, and weight management professionals for effectively evaluating and dealing with the looming crisis. The most effective way to curtail development of this problem is to prevent expansion of the obesity pandemic in both children and adults. This is a critical challenge since there are already 110 million children and adolescents and 640 million adults with obesity worldwide who constitute the at-risk
pool for development of obesity-accelerated malignancies (94,95). Moreover, this at-risk population is even further expanded by the demonstration that the effects of overweight/obesity may have a long latent period and, in some cases, precede the diagnosis of malignancy by decades. Thus, an important goal for medical professionals and supporting agencies is to encourage obesity prevention, weight loss, and increased physical activity in both children and young adults (3). In some cases, extreme measures such as bariatric surgery are being considered in children and young adults with obesity because of the potential consequences of disorders such as polycystic ovary syndrome and fatty liver disease. Special attention needs to be focused on detecting, monitoring, and reversing metabolic syndrome in all patients, especially young adults.

Interestingly, the demonstration that ASA reduced incidence of CRC in young HNPCC patients with obesity (51) indicates the importance of further research to improve cancer prevention strategies in patients with hereditary cancer syndromes and more broadly in young adults with overweight or obesity (51). Like ASA, potential chemopreventive agents for young adults with obesity must be relatively nontoxic and safe for long-term administration. Metformin, extensively used for treatment of type 2 diabetes mellitus, has already been shown, at low dose, to be safe and reduce recurrence of colorectal adenoma in nondiabetic patients following initial polyp resection (96). Because obesity-promoted cancers have been shown in some cases to involve epigenetic changes (97), other potential opportunities for chemoprevention in young adults include the use of epigenetic targeted therapies (98).

In terms of screening the young adult population for early signs of malignancy, what is clearly needed is a series of easily administered, minimally invasive, and cost-effective screening tools. These might include training and encouraging young women with obesity to perform breast self-examination, regular thyroid palpation by medical and dental practitioners, stool DNA testing for both upper and lower gastrointestinal pathologies (99), and further development of screening blood tests for circulating DNA, circulating tumor cells, and other potential biomarkers (100). Because overweight and obesity are lifestyle consequences, it is possible that they can be sufficiently altered by lifestyle modifications to avert the impending expansion of young adult cancers.

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References
1. Bleyer A, Barr R, Hayes-Lattin B, Thomas D, Ellis C, Anderson B. The distinctive biology of cancer in adolescents and young adults. Nat Rev Cancer 2008;8:288-298.
2. Apovian CM. The obesity epidemic—understanding the disease and the treatment. N Engl J Med 2016;374:177-179.
3. Dietz WH. Obesity and excessive weight gain in young adults: new targets for prevention. JAMA 2017;318:241-242.
4. Calle EE, Kaaks R. Overweight, obesity and cancer: epidemiological evidence and proposed mechanisms. Nat Rev Cancer 2004;4:579-591.
5. Berger NA. Obesity and cancer pathogenesis. Ann N Y Acad Sci 2014;1311:57-76.
6. Lauby-Secretan B, Scoccianti C, Loomis D, Grosse Y, Bianchini F, Straif K. Body fatness and cancer—viewpoint of the IARC Working Group. N Engl J Med 2016;375:794-798.
7. Cleary MP. Impact of obesity on development and progression of mammary tumors in preclinical models of breast cancer. J Mammary Gland Biol Neoplasia 2013;18:333-343.
8. Doernier SK, Reis ES, Leung ES, et al. High-fat diet-induced complement activation mediates intestinal inflammation and neoplasia, independent of obesity. Mol Cancer 2016;14:953-965.
9. Dawson DW, Hertzler K, Moro A, et al. High-fat, high-calorie diet promotes early pancreatic neoplasia in the conditional KrasG12D mouse model. Cancer Prev Res (Phila) 2013;6:1064-1073.
10. Khasawneh J, Schulz MD, Walsh A, et al. Inflammation and mitochondrial fatty acid beta-oxidation link obesity to early tumor promotion. Proc Natl Acad Sci U S A 2009;106:3354-3359.
11. Hill-Baskin AE, Markiewski MM, Buchner DA, et al. Diet-induced hepatocellular carcinoma in genetically predisposed mice. Hum Mol Genet 2009;18:2975-2988.
12. Nakagawa H. Recent advances in mice models of obesity- and nonalcoholic steatohepatitis-associated hepatocarcinogenesis. World J Hepatol 2015;7:2110-2118.
13. Kim T-M, Laird PW, Park PJ. The landscape of microsatellite instability in colorectal and endometrial cancer genomes. Cell 2013;155:858-868.
14. Lwin ST, Olechowicz SW, Fowler JA, Edwards CM. Diet-induced obesity promotes a myeloma-like condition in vivo. Leukemia 2015;29:507-510.
15. Liu Y, Metzinger MN, Lewellen KA, et al. Obesity contributes to ovarian cancer metastatic success through increased lipogenesis, enhanced vascularity, and decreased infiltration of m1 macrophages. Cancer Res 2015;75:5046-5057.
16. Yu W, Cline M, Maxwell LG, et al. Dietary vitamin D exposure prevents obesity-induced increase in endometrial cancer in Pten−/+ mice. Cancer Prev Res (Phila) 2015;8:1246-1258.
17. Quante M, Bhagat G, Abrams JA, et al. Bile acid and inflammation activate gastric cardi stem cells in a mouse model of Barrett-like metaplasia. Cancer Cell 2012;21:36-51.
18. Bleyer A, Barr R. Cancer in young adults 20 to 39 years of age: overview. Semin Oncol 2009;36:194-206.
19. American Cancer Society. Cancer Facts and Figures 2017. Atlanta, Georgia: American Cancer Society; 2017.
20. Howlender N, Noone A, Krapcho M, et al., eds. SEER Cancer Statistics Review, 1975-2014. Bethesda, MD: National Cancer Institute; 2016. https://seer.cancer.gov/csr/1975_2014/. Published April 2017. Updated June 28, 2017. Accessed September 1, 2017.
21. Arnold M, Pandeya N, Byrnes G, et al. Global burden of cancer attributable to high body-mass index in 2012: a population-based study. Lancet Oncol 2015;16:35-46.
22. Kinzler KW, Vogelstein B. Lessons from hereditary colorectal cancer. Cell 1996;87:159-170.
23. Jones S, Chen WD, Parmigiani G, et al. Comparative lesion sequencing provides insights into tumor evolution. Proc Natl Acad Sci U S A 2008;105:4283-4288.
24. Wood LD, Parsons DW, Jones S, et al. The genomic landscapes of human breast and colorectal cancers. Science 2007;318:1108-1113.
25. Cohen AJ, Saikahova A, Corradin O, et al. Hotspots of aberrant enhancer activity punctuate the colorectal cancer epigenome. Nat Commun 2017;8:14400. doi: 10.1038/ncomms14400.
26. Cohen DH, LeRoith D. Obesity, type 2 diabetes, and cancer: the insulin and IGF connection. Endocr Relat Cancer 2012;19:F27-F45.
27. Karuwanathi S, Levi L, Devicchio J, et al. RBP4-STRAX pathway drives cancer stem cell maintenance and mediates high-fat diet-induced colon carcinogenesis. Stem Cell Reports 2017;9:438-450.
28. Iyengar NM, Brown KA, Zhou XK, et al. Metabolic obesity, adipose inflammation and elevated breast aromatase in women with normal body mass index. Cancer Prev Res (Phila) 2017;10:235-243.
29. Chang SH, Luo S, Thomas TS, et al. Obesity and the transformation of monoclonal gammapathy of undetermined significance to multiple myeloma: a population-based cohort study. J Natl Cancer Inst 2017;109. doi: 10.1093/jnci/djw264.
30. Kim EI, Choi M-R, Park H, et al. Dietary fat increases solid tumor growth and metastasis of 4T1 murine mammary carcinoma cells and mortality in obesity-resistant BALB/c mice. Breast Cancer Res 2011;13:R78. doi:10.1186/bcr2927.
31. Novosyadlyy R, Leroith D. Insulin-like growth factors and insulin: at the crossroad between tumor development and longevity. J Gerontol A Biol Sci Med Sci. 2012;67:640-651.
32. Zheng Q, Dunlap SM, Zhu J, et al. Leptin deficiency suppresses MMTV-Wnt-1 mammary tumor growth in obese mice and abrogates tumor initiating cell survival. Endocr Relat Cancer 2011;18:491-503.
33. O’Connell JB, Maggard MA, Liu JH, Etzioni DA, Livingston EH, Ko CY. Rates of colon and rectal cancers are increasing in young adults. Am Surg 2003;69:866-872.
34. Connell LC, Mota JM, Braghiroli MI, Hoff PM. The rising incidence of younger patients with colorectal cancer: questions about screening, biology, and treatment. Curr Treat Options Oncol 2017;18:23. doi:10.1007/s11864-017-0463-3.
35. Siegel RL, Jemal A, Ward EM. Increase in incidence of colorectal cancer among young men and women in the United States. Cancer Epidemiol Biomarkers Prev 2009;18:1695-1698.

36. Grahn SW, Varma MG. Factors that increase risk of colon polyps. Clin Colon Rectal Surg 2009;22:247-255.

37. Atkin W, Wooldrage K, Brenner A, et al. Adenoma surveillance and colorectal cancer in young adults. J Clin Oncol 2015;33:3544-3549.

38. Kim JY, Jung YS, Park JH, et al. Different risk factors for advanced colorectal neoplasm in young adults. World J Gastroenterol 2016;22:3611-3620.

39. Kotsopoulos J, Olopado OI, Ghadirian P, et al. Changes in body weight and the risk of breast cancer in BRCA1 and BRCA2 mutation carriers. Ann Intern Med 2011;154:14-22.

40. You YN, Xing Y, Feig BW, Chang GJ, Cormier JN. Young-onset colorectal cancer: is it time to pay attention? Arch Intern Med 2012;172:287-289.

41. Bailey CE, Hu CY, You YN, et al. Increasing disparities in the age-related incidences of colon and rectal cancers in the United States, 1975-2010. JAMA Surg 2015;150:17-22.

42. Siegel RL, Miller KD, Fedewa SA, et al. Colorectal cancer statistics, 2017. CA Cancer J Clin 2017;67:177-193.

43. Le Marchand L, Wilkins LR, Mi MP. Obesity in youth and middle age and risk of colorectal cancer. Cancer Causes Control 1999;10:349-354.

44. Ortiz AP, Thompson CL, Chak A, Berger NA, Li L. Insulin resistance, central obesity, and risk of colorectal adenomas. Cancer 2012;118:1774-1781.

45. Atkin W, Wooldrige K, Brenner A, et al. Adenoma surveillance and colorectal cancer incidence: a retrospective, multicentre, cohort study. Lancet Oncol 2017;18:823-834.

46. Grahn SW, Varma MG. Factors that increase risk of colon polyps. Clin Colon Rectal Surg 2009;22:247-255.

47. Haggar FA, Boushey RP. Colorectal cancer epidemiology: incidence, mortality, survival, and risk factors. Clin Colon Rectal Surg 2009;22:191-197.

48. Hahn MM, de Voer RM, Hoogerbrugge N, Ligtenberg MJ, Kuiper RP, van Kessel AG. The genetic heterogeneity of colorectal cancer predisposition - guidelines for gene discovery. Curr Opin Genet Dev 2005;15:349-356.

49. van Duijnhoven FJ, Botma A, Winkels R, Nagengast FM, Vasen HF, Kampman E. Do lifestyle factors influence colorectal cancer risk in Lynch syndrome? Fam Cancer 2013;12:285-293.

50. Mork ME, You YN, Jing J, et al. High prevalence of hereditary cancer syndromes in adolescents and young adults with colorectal cancer. J Clin Oncol 2015;33:3546-3549.

51. Movahedi M, Bishop DT, Macrae F, et al. Obesity, aspirin, and risk of colorectal cancer in carriers of hereditary colorectal cancer: a prospective investigation in the CAPP2 Study. J Clin Oncol 2015;33:3591-3597.

52. Anders CK, Johnson R, Litton J, Phillips M, Bleyer A. Breast cancer before age 40. Semin Oncol 2016;43:359-368.

53. Kim SE, Shim KN, Jung SA, et al. An association between obesity and the prevalence of colonic adenoma according to age and gender. J Gastroenterol Hepatol 2007;22:616-623.

54. Levi Z, Kark JD, Barchana M, et al. Measured body mass index in adolescence and the incidence of colorectal cancer in a cohort of 1.1 million males. Cancer Epidemiol Biomarkers Prev 2011;20:2524-2531.

55. Kim JY, Jung YS, Park JH, et al. Different risk factors for advanced colorectal neoplasm in young adults. World J Gastroenterol 2016;22:3611-3620.

56. Kotsopoulos J, Olopado OI, Ghadirian P, et al. Changes in body weight and the risk of breast cancer in BRCA1 and BRCA2 mutation carriers. Ann Intern Med 2011;154:14-22.
93. Castro FA, Koshol J, Hsing AW, Devesa SS. Biliary tract cancer incidence in the United States—Demographic and temporal variations by anatomic site. *Int J Cancer* 2013;133:1664-1671.

94. Ezzati MN-RC. Trends in adult body-mass index in 200 countries from 1975 to 2014: a pooled analysis of 1698 population-based measurement studies with 19.2 million participants. *Lancet* 2016;387:1377-1396.

95. Afshin A, Forouzanfar MH, Reitsma MB, et al. Health effects of overweight and obesity in 195 countries over 25 years. *N Engl J Med* 2017;377:13-27.

96. Higurashi T, Hosono K, Takahashi H, et al. Metformin for chemoprevention of metachronous colorectal adenoma or polyps in post-polypectomy patients without diabetes: a multicentre double-blind, placebo-controlled, randomised phase 3 trial. *Lancet Oncol* 2016;17:475-483.

97. Rossi EL, de Angel RE, Bowers LW, et al. Obesity-associated alterations in inflammation, epigenetics, and mammary tumor growth persist in formerly obese mice. *Cancer Prev Res (Phila)* 2016;9:339-348.

98. Jones PA, Isa J, Baylin S. Targeting the cancer epigenome for therapy. *Nat Rev Genet* 2016;17:630-641.

99. Moinova H, Leidner RS, Ravi L, et al. Aberrant vimentin methylation is characteristic of upper gastrointestinal pathologies. *Cancer Epidemiol Biomarkers Prev* 2012;21:594-600.

100. Friedrich MJ. Going with the flow: the promise and challenge of liquid biopsies. *JAMA* 2017;318:1095-1097.