Routine Clinical Measures of Adiposity as Predictors of Visceral Fat in Adolescence: A Population-Based Magnetic Resonance Imaging Study

Katie Goodwin¹, Catriona Syme¹, Michal Abrahamowicz², Gabriel T. Leonard³, Louis Richer⁴, Michel Perron⁵, Suzanne Veillette⁶, Daniel Gaudet⁷, Tomas Paus⁵, Zdenka Pausova*⁷

¹ Hospital for Sick Children, University of Toronto, Toronto, Canada, ² Department of Epidemiology, Biostatistics and Occupational Health, McGill University, Montreal, Canada, ³ Montreal Neurological Institute, McGill University, Montreal, Canada, ⁴ Department of Psychology, Université du Québec à Chicoutimi, Chicoutimi, Canada, ⁵ Department of Human Sciences, Université du Québec à Chicoutimi, Chicoutimi, Canada, ⁶ Community Genomic Centre, Université de Montréal, Chicoutimi Hospital, Chicoutimi, Canada, ⁷ Rotman Research Institute, University of Toronto, Toronto, Canada

Abstract

Objective: Visceral fat (VF) increases cardiometabolic risk more than fat stored subcutaneously. Here, we investigated how well routine clinical measures of adiposity, namely body mass index (BMI) and waist circumference (waist), predict VF and subcutaneous fat (SF) in a large population-based sample of adolescents. As body-fat distribution differs between males and females, we performed these analyses separately in each sex.

Design and Methods: VF and SF were measured by magnetic resonance imaging in 1,002 adolescents (482 males, age 12–18 years). Relationships of BMI and waist with VF and SF were tested in multivariable analyses, which adjusted for potentially confounding effects of age and height.

Results: In both males and females, BMI and waist were highly correlated with VF and SF, and explained 55–76% of their total variance. When VF was adjusted for SF, however, BMI and waist explained, respectively, only 0% and 4% of VF variance in males, and 4% and 11% of VF variance in females. In contrast, when SF was adjusted for VF, BMI and waist explained, respectively, 36% and 21% of SF variance in males, and 48% and 23% of SF variance in females. These relationships were similar during early and late puberty.

Conclusions and Relevance: During adolescence, routine clinical measures of adiposity predict well SF but not VF. This holds for both sexes and throughout puberty. Further longitudinal studies are required to assess how well these measures predict changes of VF and SF over time. Given the clinical importance of VF, development of cost-effective imaging techniques and/or robust biomarkers of VF accumulation that would be suitable in everyday clinical practice is warranted.

Introduction

Obesity is a major public health problem [1,2]. Due to its currently high prevalence and adverse effects on health [1,2], our life expectancy has been projected to decline for the first time since the Great Depression [3]. This is in part due to obesity increasing significantly risk for cardio-metabolic diseases (CMD), such as cardiovascular disease and type-2 diabetes mellitus [2,4], which in turn are the major causes of morbidity and mortality in the industrialized world. This obesity-related risk for CMD is mainly the consequence of adipose tissue releasing a number of adipocytokines that, once released into the circulation, promote the development of CMD [5].

The relationship between obesity and CMD is not simple, however. About 30% of adult individuals who are classified as “obese” are cardiometabolically healthy and over 20% who are classified as being of “normal weight” are cardiometabolically abnormal [6]. To some extent, this may be an artefact of the measurement method. For over 150 years, obesity has been
assessed with body mass index (BMI, weight/height^2), which is an imprecise, and possibly misleading [7], metric of body fat. This is because BMI (and waist circumference [waist]) are influenced not only by fat mass but also by muscle mass and bone mass, among others [8].

Further, obesity-related risk for CMD increases not only with the quantity but also with a specific distribution of body fat – individuals who store body fat viscerally rather than elsewhere in the body (mostly subcutaneously) are at a greater risk for CMD [9–11]. This relationship is seen not only in adults, but also in children and adolescents [12–16]. Several mechanistic pathways have been proposed to underlie the link between visceral fat (VF) and CMD - VF, as compared with subcutaneous fat (SF), exhibits a more adverse secretory profile and higher lipid turnover [4,17–19]. Further, VF but not SF drains directly to the portal circulation and liver, where it enhances dyslipidemia and insulin resistance, key mediators of the link between obesity and CMD [4,20,21]. Consistent with these differences between VF and SF, it has been demonstrated that surgical removal of VF but not that of SF improves cardio-metabolic health in humans and experimental animals [22–24]. Moreover, a growing body of research suggests that SF may even be cardiometabolically protective in overweight and obesity [14,25,26]. This effect has been related to the so-called “expandability” of SF, which is thought to be a biological property of SF that allows excess body-fat to accumulate preferentially in this fat depot and thus protects the body from the cardiometabolically adverse accumulation of VF [27,28].

Thus, given the biological and clinical differences between VF and SF and hence the importance of quantifying them separately, the aim of the present study was to investigate how well routine clinical measures of adiposity, namely BMI and waist, predict VF and SF measured directly with magnetic resonance imaging (MRI) in a large population-based sample of adolescents (n=1,002). We focused here on adolescence, as it is an understudied period of human development [29] during which the initial stages of CMD may emerge [13,30–34].

**Methods**

**Adolescent sample**

The study sample consisted of White Caucasian males (n=482) and females (n=520), aged 12 to 18 years, who were recruited via high schools from the Saguenay-Lac St. Jean region of Quebec, Canada, as part of the Saguenay Youth Study (SYS) [35]. The SYS is a population-based cross-sectional study of cardio-metabolic and mental health during adolescence. Assent of the adolescents and written consent of the parents were obtained before data collection, and the Research Ethics Committees of the Chicoutimi Hospital (Chicoutimi, Canada) and the Hospital for Sick Children (Toronto, Canada) approved the study. The current sample of 1,002 adolescents was recruited and tested between November 2003 and February 2012.

**Routine clinical measures of adiposity**

Height (0.1-cm precision), weight (0.1-kg precision), waist (0.1-cm precision), hip circumference (0.1-cm precision), and suprailiac skinfold-thickness (suprailiac skinfold, 1-mm precision) were measured by trained staff using standard operating procedures [35]. Waist was measured at the level of the natural waist, which is the narrowest part of the torso, as seen from the anterior aspect. Hip circumference was measured at the level of the maximum extension of the buttocks. Suprailiac skinfold was measured in the midaxillary line immediately superior to the iliac crest. BMI was calculated as weight (in kg) divided by height (in m)^2. Waist to hip ratio (waist/hip) was calculated as waist (in cm) divided by hip circumference (in cm).

**Magnetic resonance imaging of abdominal VF and SF**

VF and SF were quantified with MRI from axial, 10-mm thick (with in-plane resolution 1.56 x 1.56 mm^2), heavily T1-weighted, spin-echo (TR/TE = 200 ms/20 ms) scans, which were taken along the abdomen with a Phillips 1.0-T magnetic resonance scanner. A single slice at the level of the umbilicus was used to quantify VF and SF. The single slice images were smoothed using an adaptive bilateral filter to remove image noise while preserving edge information. A standard region growing algorithm written in MatLab (R2011a, MathWorks, Natick, MA) was used to obtain an initial fat classification map. An iterative refinement procedure corrected false positives and negatives using morphological operators, including hysteresis, thresholding over small neighbourhoods, and median filtering to remove salt-and-pepper noise. The resultant image was manually segmented into SF and VF using Adobe Photoshopp (CS5, Adobe Systems Incorporated, San Jose, CA). SF was defined as the area of adipose tissue between the skin and the outer aspect of abdominal musculature. VF was defined as the as the area of adipose tissue within the inner aspect of abdominal musculature and outside of abdominal organs. A histogram counting algorithm written in MatLab (R2011a, MathWorks, Natick, MA) computed the total number of pixels for each type of fat (VF and SF). All 1,002 scans were segmented by a single individual (KG). This segmentation was validated by another individual (CS) in 586 scans (VF: r=0.92, SF: r=0.99).

**Pubertal development**

Puberty stage (1 to 5) was assessed using Puberty Development Scale, which is an 8-item self-report measure of physical development based on Tanner stages [36]. With this tool, separate forms for males and females are used and the 5 puberty stages (i.e., 1. pre-pubertal, 2. beginning pubertal, 3. mid-pubertal, 4. advanced pubertal and 5. post-pubertal) are assessed based on answers to questions regarding pubic hair, growth in stature, menarche for girls and voice changes in males. This tool was validated against physician’s assessment of pubertal development [37] and correlates well with plasma level of sex hormones [38].
Statistical methods

Descriptive statistics used to characterize the study population included means and standard deviations. Our main analyses were aimed at estimating the proportion of variance shared between each of two main clinical measures of adiposity, namely BMI and waist, and VF or SF. We ran two sets of multivariable linear regression models, that all adjusted for inter-individual differences in age and height, and used either VF or SF as the continuous dependent variable. In the first sets of analyses, the models that predicted VF did not adjust for SF, and vice versa; whereas in the second sets of analyses, they did. The latter sets of analyses were intended to assess if, and to what extent, BMI and waist predict VF even among adolescents with the same SF, and vice versa. The same analytical methods were also employed to examine if VF and SF can be predicted by suprailiac skinfold and waist/hip, which are additional anthropometric measures of central adiposity that could be used in clinical setting. Similar to other studies [29], marked sex differences in body-fat distribution were evident in our sample (Table 1). Therefore, we run all our analyses separately in males and females.

As secondary analyses, we examined whether pubertal development influences the observed relationships. To this end, we added the two-way interactions of puberty stage (dichotomized as early [stages 1-3] vs. late [stages 4-5] puberty) with each of the tested main independent variables (BMI, waist, suprailiac skinfold and waist/hip). Again, these analyses were carried out in each sex separately.

In preliminary analyses, distributions of all continuous variables were assessed for the normality assumption on which the statistical inference about the linear model estimates relies. The values of variables for which the empirical distribution showed substantial positive skeweness (normality tested with K–S statistics), namely BMI, waist, suprailiac skinfold, waist/hip, VF and SF were all log transformed, using logarithm with base 10, which improved the fit of the multivariable models in the main analyses (data not shown).

All statistical analyses were performed using JMP software (Release 9, SAS Institute Inc., Cary, NC).

Results

Characteristics of studied adolescent males and females

Mean ages of males (n=482) and females (n=520) differed by only 1 month, but, as expected [39], males were at an earlier stage of pubertal development (p<0.0001, Table 1). Males and females had similar BMI (p=0.75) and VF (p=0.17), but males (vs. females) showed higher waist (p<0.0001) and waist/hip (p<0.0001), and lower suprailiac skinfold (p<0.0001) and SF (p<0.0001, Table 1).

As expected, SF was closely associated with VF; the two measures shared 77% (p<0.0001) and 64% (p<0.0001) of variance in males and females, respectively (Figure 1). The slope of this relationship was steeper in males than females (p=0.04 for the test of interaction with sex) indicating that, for a given quantity of SF, males compared with females have more VF (Figure 1).

Routine clinical measures of adiposity as predictors of abdominal VF and SF

BMI was closely associated with VF and SF (p<0.0001) in both sexes; it explained 58% and 55% of variance of VF and 72% and 76% of variance of SF in males and females of the same age and height, respectively. Relationships of waist with VF and SF were equally strong (p<0.0001); it explained 61% (p<0.0001) and 56% (p<0.0001) of variance of VF and 67% (p<0.0001) and 62% (p<0.0001) of variance of SF in males and females of the same age and height, respectively (Figure S1 in File S1).

Table 1. Basic characteristics and adiposity measures of studied adolescent males and females.

| Variables                      | Males Mean ±SD | Females Mean ±SD | p-value |
|--------------------------------|----------------|------------------|---------|
| Number                         | 482            | 518              |         |
| Age (months)                   | 180 ±21        | 181 ±23          | 0.18    |
| Height (cm)                    | 167 ±11        | 160 ±6.7         | <0.0001 |
| Puberty stage (1-5)            | 10/61/184/197/29 | 3/37/25/29/143  | <0.0001 |
| Stage 1                        | 2%             | 1%               |         |
| Stage 2                        | 13%            | 1%               |         |
| Stage 3                        | 38%            | 14%              |         |
| Stage 4                        | 41%            | 57%              |         |
| Stage 5                        | 6%             | 27%              |         |
| Puberty stage (Early/Late)     | 255/226        | 83/439           | <0.0001 |
| Body mass index                |                |                  |         |
| kg/m²                          | 21.7 ±4.3      | 21.7 ±4.0        | 0.87    |
| log kg/m²                      | 1.33 ±0.08     | 1.33 ±0.06       | 0.75    |
| percentile                     | 58.8 ±28.8     | 56.1 ±27.3       | 0.14    |
| Waist                          |                |                  |         |
| cm                             | 75.1 ±10       | 71.0 ±8.8        | <0.0001 |
| log cm                         | 1.87 ±0.06     | 1.85 ±0.05       | <0.0001 |
| Hip                            |                |                  |         |
| cm                             | 88.9 ±10       | 89.9 ±9.9        | 0.11    |
| log cm                         | 1.95 ±0.05     | 1.95 ±0.05       | 0.08    |
| Suprailiac skinfold            |                |                  |         |
| mm                             | 16.8 ±12       | 20.6 ±10         | <0.0001 |
| log mm                         | 1.13 ±0.28     | 1.26 ±0.21       | <0.0001 |
| Waist/hip                      |                |                  |         |
| cm/cm                          | 0.848 ±0.056   | 0.791 ±0.062     | <0.0001 |
| log cm/cm                      | -0.073 ±0.028  | -0.103 ±0.033    | <0.0001 |
| Waist/height                   |                |                  |         |
| cm/cm                          | 0.440 ±0.084   | 0.439 ±0.071     | 0.88    |
| log cm/cm                      | -0.350 ±0.051  | -0.355 ±0.050    | 0.14    |
| Visceral fat                   |                |                  |         |
| cm²                            | 230.49 ±224.98 | 208.78 ±141.99   | 0.07    |
| log cm²                        | 2.22 ±0.34     | 2.24 ±0.25       | 0.17    |
| Subcutaneous fat               |                |                  |         |
| cm²                            | 1058.57 ±1028.26 | 1450.07 ±963.33 | <0.0001 |
| log cm²                        | 2.86 ±0.36     | 3.08 ±0.27       | <0.0001 |

Unadjusted mean ± standard deviation for raw and log-transformed values of relevant characteristics are shown for the studied males and females. P values indicate statistical significance of differences between males and females evaluated with 2-sided t test.

doi: 10.1371/journal.pone.0079896

Clinical Measures of Visceral Fat in Adolescence

PLOS ONE | www.plosone.org

November 2013 | Volume 8 | Issue 11 | e79896
However, in the models predicting VF adjusted for SF, BMI and waist explained, respectively, only 0% (p=0.69) and 4% (p<0.0001) of VF variance (in males); similarly in females, BMI and waist explained, respectively, just 4% (p<0.0001) and 11% (p<0.0001) of VF variance (Figure 2). In contrast, in the models predicting SF adjusted for VF, BMI and waist explained, respectively, substantial proportions of SF variance - 36% (p<0.0001) and 21% (p<0.0001) in males and 48% (p<0.0001) and 23% (p<0.0001) in females (Figure 2). In other words, among both males and females with the same SF, BMI and waist did not relate to VF but, among adolescents with the same VF, these two measures did relate well to SF.

Major changes in body size and composition happen during puberty, with accelerated weight gain occurring in most boys...
and girls at stages 4 and 5 of pubertal development [40]. Therefore, we examined whether the relationships of BMI and waist circumference with each VF and SF (while adjusting for each other) are shown in adolescent males and females. All relationships were also adjusted for potentially confounding effects of age and height when appropriate.

doi: 10.1371/journal.pone.0079896.g002

and girls at stages 4 and 5 of pubertal development [40]. Therefore, we examined whether the relationships of BMI and waist circumference with eachVF and SF (while adjusting for each other) are shown in adolescent males and females. All relationships were also adjusted for potentially confounding effects of age and height when appropriate.

Additional anthropometric measures as predictors of abdominal VF and SF

In addition to BMI and waist, we investigated two other anthropometric measures of adiposity that could be used in clinics as predictors of VF and SF, namely suprailiac skinfold and waist/hip. Suprailiac skinfold showed strong relationships with VF and SF in both sexes, explaining 62% (p<0.0001) and 52% (p<0.0001) of VF variance and 72% (p<0.0001) and 64% (p<0.0001) of SF variance in males and females, respectively (Figure S2 in File S1). But again, when VF was adjusted for SF, suprailiac skinfold explained only 2% (p=0.002) and 5% (<0.001) of VF variance in males and females, respectively, and when SF was adjusted for VF, it explained 27% (p<0.0001) and 26% (p<0.0001) of SF variance in males and females, respectively (Figure S3 in File S1). As such, suprailiac skinfold compared with BMI and waist showed a similar potential to predict VF and SF.

Waist/hip, however, demonstrated a different pattern of relationships. It showed substantially weaker relationships with both VF or SF (as compared with BMI, waist and suprailiac skinfold) – it explained only 22% (p<0.0001) and 20% (p<0.0001) of VF variance and 12% (p<0.0001) and 6% (p<0.0001) of SF variance in males and females, respectively (Figure S2 in File S1). Further, when VF was adjusted for SF, waist/hip explained 5% (p<0.0001) and 6% (p<0.0001) of VF variance in males and females, respectively, which was similar to the other anthropometric measures, but unlike them, waist/hip explained 0% of variance in both sexes (p=0.81 and 0.33, respectively) when SF was adjusted for VF (Figure S3 in File S1). As such, waist/hip compared with BMI, waist and suprailiac skinfold showed a similarly low potential to predict VF and an even lower potential to predict SF.

The relationships of both these additional anthropometric measures of adiposity (i.e., suprailiac skinfold and waist/hip) with VF and SF did not differ between early and late puberty groups (p>0.29 and p>0.12 for the tests of interaction with puberty stage in males and females, respectively).

Discussion

The results of the present study demonstrate that, during adolescence, routine clinical measures of adiposity, namely BMI and waist, predict SF but not VF. These relationships are similar in both sexes and exist throughout puberty.

The current study is a large-scale investigation of adolescents (n=1,002) testing BMI and waist as predictors of VF and SF measured directly with MRI. A few such (>500 individuals) investigations have been conducted in adults [41–43] but none in children or adolescents. In the latter age category (children or adolescents), only smaller-scale studies have been performed [44–46]. Irrespective of their size or participants’ age, all of these studies have shown, similarly to the current one (Figure S1 in File S1), that BMI and waist are closely related to both VF and SF and explain large proportions of their respective variances (BMI: 37-69% of VF and 74-89% of SF; waist: 53-70% of VF and 67-86% of SF) [41–46]. But because VF and SF are closely associated with each other (64-77%, Figure 1), it is likely these relationships are driven mainly by global adiposity. Therefore, in the present study (in contrast to previous studies [41–45]), we examined how BMI and waist relate to VF and SF independently of each other – thus, we examined whether BMI and waist can predict VF- and SF-specific quantities of body fat. These results showed that BMI and waist explain a very small proportion of VF-specific variance (0-11%, Figure 2) but a substantially larger proportion of SF-specific variance (21-48%, Figure 2), indicating that both BMI and waist predict mainly SF but not VF.
In the present study, we examined two additional anthropometric measures of adiposity that could be used in clinics as predictors of VF and SF – these were suprailiac skinfold and waist/hip. Both measures performed either similarly or even worse than BMI and waist (Figures S2 and S3 in File S1).

The current study has potential limitations. First, it was carried out in White Caucasians and its findings may not be generalizable to other ethnicities. Second, we quantified VF and SF using a single slice. Single-slice data may not be as accurate as multi-slice data in quantifying VF volume [47], but they have been deemed to be a reasonable approximation [48–50]. Third, we quantified VF and SF at the level of the umbilicus. This level may be problematic, as it may correspond to varying locations of the lumbar vertebrae [48,51]. In the present study, we observed that the umbilicus was located within a segment of the body that is 5 to 10 cm above the L4/L5 vertebral disk in most our participants (20 out of 20 randomly selected individuals). VF assessed as a slice area within this segment has been reported as most closely associated with VF volume [48,49], as well as with its age-related increases [52] and risk for CMD [53].

The results of the present large-scale study suggest that, in adolescence, BMI and waist circumference cannot be used to estimate VF. This poses a challenge for the practice of personalized (preventive) medicine, as youth at risk for obesity-related CMD cannot be easily identified using these clinical measures [54]. Furthermore, the use of BMI and waist circumference may obscure our understanding of obesity-related CMD and thus development of new effective treatments. As described in Introduction, VF vs. SF is more closely associated with obesity-related risk for CMD [9–16]. This is because VF vs. SF is more closely located to the portal circulation [4,20,21] and demonstrates a more adverse secretory profile [4,17–19]. Further, SF may even be cardiometabolically protective in overweight and obesity [14,25,26] due to its “expandability” and thus protection from the cardiometabolically adverse accumulation of VF [27,28].

Given the clear limitation of currently used clinical measures of adiposity (BMI and waist circumference) to predict VF, development of cost-effective imaging techniques to quantify VF that would be suitable for everyday clinical practice and/or identification of robust circulating biomarkers of VF accumulation are warranted.

Supporting Information

**File S1.** Supporting table and figures. Table S1, Basic characteristics and adiposity measures for early and late puberty males and females. Figure S1, BMI and waist circumference as predictors of VF and SF quantities (VF and SF not adjusted for each other). Figure S2, suprailiac skinfold thickness and waist-to-hip ratio as predictors of VF and SF quantities (VF and SF not adjusted for each other). Figure S3, suprailiac skinfold thickness and waist-to-hip ratio as predictors of VF and SF quantities (VF and SF adjusted for each other).

(DOC)

Acknowledgements

We thank the following individuals for their contributions in acquiring data: Manon Bernard (database architect, The Hospital for Sick Children), Helene Simard and her team of research assistants (Cégep de Jonquière) and Dr. Rosanne Aleong (program manager, Rotman Research Institute). Dr. Zdenka Pausova had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. The data are deposited in The Saguenay Youth Study database and are accessible via request to senior investigators of the study (ZP and TP).

Author Contributions

Conceived and designed the experiments: ZP TP KG MA LR. Performed the experiments: TP MA KG. Analyzed the data: CS KG MA TP. Wrote the manuscript: ZP MA KG. Editing manuscript: TP MA.

References

1. Anis AH, Zhang W, Bansback N, Guh DP, Amarsi Z et al. (2010) Obesity and overweight in Canada: an updated cost-of-illness study. Obes Rev 11: 31-40. doi:10.1111/j.1467-789X.2009.00579.x. PubMed: 19413707.

2. Must A, Spadano J, Coakley EH, Field AE, Colditz G et al. (1999) The disease burden associated with overweight and obesity. JAMA 282: 1523-1529. doi:10.1001/jama.282.16.1523. PubMed: 10546891.

3. Olshansky SJ, Passaro DJ, Hershow RC, Layden J, Carnes BA et al. (2005) A potential decline in life expectancy in the United States in the 21st century. N Engl J Med 352: 1138-1145. doi:10.1056/NEJMsr043743. PubMed: 15784668.

4. Cornier MA, Després JP, Davis N, Grossniklaus DA, Klein S et al. (2011) Assessing adiposity: a scientific statement from the American Heart Association. Circulation 124: 1996-2019. doi:10.1161/CIR.0b013e318233bc5a. PubMed: 21947291.

5. Pausova Z (2006) From big fat cells to high blood pressure: a pathway to obesity-associated hypertension. Curr Opin Nephrol Hypertens 15: 173-178. doi:10.1097/01.mnh.0000214775.42103.a5. PubMed: 16481885.

6. Wildman RP, Muntner P, Reynolds K, McGinn AP, Rajpathak S et al. (2006) The obese without cardiometabolic risk factor clustering and the normal weight with cardiometabolic risk factor clustering: Prevalence and correlates of 2 phenotypes among the us population (nhanes 1999-2004). Arch Intern Med 168: 1617-1624. doi:10.1001/archinte.168.15.1617. PubMed: 18695075.

7. Pou KM, Massaro JM, Hoffmann U, Lieb K, Vasan RS et al. (2009) Patterns of abdominal fat distribution: the Framingham Heart Study. Diabetes Care 32: 481-485. doi:10.2337/dc08-1359. PubMed: 19074995.

8. Müller MJ, Lagerpuch M, Enderle J, Schultz B, Heller M et al. (2012) Beyond the body mass index: tracking body composition in the pathogenesis of obesity and the metabolic syndrome. Obes Rev 13 Suppl 2: 6-13. doi:10.1111/j.1467-789X.2012.01033.x. PubMed: 23107255.

9. Fox CS, Massaro JM, Hoffmann U, Pou KM, Maurovich-Horvat P et al. (2007) Abdominal visceral and subcutaneous adipose tissue compartments: association with metabolic risk factors in the Framingham Heart Study. Circulation 116: 39-48. doi:10.1161/CIRCULATIONAHA.107.184046. PubMed: 17578866.

10. Demerath EW, Reed D, Rogers N, Sun SS, Lee M et al. (2008) Visceral adiposity and its anatomical distribution as predictors of the...
Clinical Measures of Visceral Fat in Adolescence

McGill HC Jr., McManus CA (2011) Starting earlier to prevent heart disease. JAMA 291: 2320-2322. doi: 10.1001/jama.291.17.2320. PubMed: 14600192.

Cook S, Wiltzmann M, Auinger P, Nguyen M, Dietz WH (2004) Prevalence of a metabolic syndrome phenotype in adolescents: findings from the third National Health and Nutrition Examination Survey, 1988-1994. Arch Pediatr Adolesc Med 157: 821-827. doi: 10.1001/archpedi.157.8.821. PubMed: 12912790.

Johnson WD, Kroon JJ, Greenway FL, Bouchard C, Ryan D et al. (2009) Prevalence of risk factors for metabolic syndrome in adolescents: National and Regional Differences. Nutr J 8: 156. doi: 10.1186/1475-2891-8-156. PubMed: 19349567.

Steinberger J, Daniels SR, Eckel RH, Hayman L, Lustig RH et al. (2009) Progress and challenges in metabolic syndrome in children and adolescents: a scientific statement from the American Heart Association Atherosclerosis, Hypertension, and Obesity in the Young Committee of the Council on Cardiovascular Disease in the Young, Council on Cardiovascular Nursing, and Council on Nutrition, Physical Activity, and Metabolism. Circulation 119: 626-647. doi: 10.1161/CIRCULATIONAHA.110.191394. PubMed: 19139390.

Taksali SE, Caprio S, Dziura J, Dufour S, Call AM et al. (2008) High visceral and low abdominal subcutaneous fat stores in the obese adolescent: a determinant of an adverse metabolic phenotype. Diabetes 57: 367-371. PubMed: 17977954.

Pausova Z, Paus T, Abrahamowicz M, Almerji J, Arbour N et al. (2007) Genes, maternal smoking, and the offspring brain and body during adolescence: design of the Saguenay Youth Study. Hum Brain Mapp 22: 502-518. doi: 10.1002/hbm.20402. PubMed: 17469173.

Peterson A, Crockett L, Richards M, Boxer A (1988) A self-report measure of pubertal status: Reliability, validity, and initial norms. J Adolescent Health 17: 117-133. doi: 10.1016/0140-058X(88)90084-9. PubMed: 3174312.

Shircliff EA, Dahl RE, Pollak SD (2009) Pubertal development: correspondence between hormonal and physical development. Child Dev 80: 327-337. doi: 10.1111/j.1465-8624.2009.01263.x. PubMed: 19711337.

Bolinder J, Kager L, Ostman J, Aner P (2003) Differences at the receptor and postreceptor levels between human omental and subcutaneous adipose tissue in the action of insulin on lipolysis. Arch Pediatr Adolesc Med 157: 1232. doi: 10.1001/archpedi.157.11.1232. PubMed: 1337993.

Björntorp P (1990) “Portali” adipose tissue as a generator of risk factors for cardiovascular disease and diabetes. Arteriosclerosis 10: 493-496. doi: 10.1161/01.ATO.10.4.493. PubMed: 2196039.

Hocking S, Samocha-Bonet D, Milner KL, Greenfield JR, Chisholm DJ (2012) Anthropometric correlates of total body fat, abdominal fat and visceral and subcutaneous adiposity in children and adolescents: National and Regional Differences. Nutr J 88: 1263-1271. doi: 10.1038/nj.2012.95. PubMed: 22710928.

30. McGill HC Jr., McManus CA (2003) Starting earlier to prevent heart disease. JAMA 289: 2320-2322. doi: 10.1001/jama.289.17.2320. PubMed: 14600192.

Cook S, Wiltzmann M, Auinger P, Nguyen M, Dietz WH (2004) Prevalence of a metabolic syndrome phenotype in adolescents: findings from the third National Health and Nutrition Examination Survey, 1988-1994. Arch Pediatr Adolesc Med 157: 821-827. doi: 10.1001/archpedi.157.8.821. PubMed: 12912790.

Johnson WD, Kroon JJ, Greenway FL, Bouchard C, Ryan D et al. (2009) Prevalence of risk factors for metabolic syndrome in adolescents: National and Regional Differences. Nutr J 8: 156. doi: 10.1186/1475-2891-8-156. PubMed: 19349567.

Steinberger J, Daniels SR, Eckel RH, Hayman L, Lustig RH et al. (2009) Progress and challenges in metabolic syndrome in children and adolescents: a scientific statement from the American Heart Association Atherosclerosis, Hypertension, and Obesity in the Young Committee of the Council on Cardiovascular Disease in the Young, Council on Cardiovascular Nursing, and Council on Nutrition, Physical Activity, and Metabolism. Circulation 119: 626-647. doi: 10.1161/CIRCULATIONAHA.110.191394. PubMed: 19139390.

Taksali SE, Caprio S, Dziura J, Dufour S, Call AM et al. (2008) High visceral and low abdominal subcutaneous fat stores in the obese adolescent: a determinant of an adverse metabolic phenotype. Diabetes 57: 367-371. PubMed: 17977954.

Pausova Z, Paus T, Abrahamowicz M, Almerji J, Arbour N et al. (2007) Genes, maternal smoking, and the offspring brain and body during adolescence: design of the Saguenay Youth Study. Hum Brain Mapp 22: 502-518. doi: 10.1002/hbm.20402. PubMed: 17469173.

Peterson A, Crockett L, Richards M, Boxer A (1988) A self-report measure of pubertal status: Reliability, validity, and initial norms. J Adolescent Health 17: 117-133. doi: 10.1016/0140-058X(88)90084-9. PubMed: 3174312.

Shircliff EA, Dahl RE, Pollak SD (2009) Pubertal development: correspondence between hormonal and physical development. Child Dev 80: 327-337. doi: 10.1111/j.1465-8624.2009.01263.x. PubMed: 19711337.

Bolinder J, Kager L, Ostman J, Aner P (2003) Differences at the receptor and postreceptor levels between human omental and subcutaneous adipose tissue in the action of insulin on lipolysis. Arch Pediatr Adolesc Med 157: 1232. doi: 10.1001/archpedi.157.11.1232. PubMed: 1337993.

Björntorp P (1990) “Portali” adipose tissue as a generator of risk factors for cardiovascular disease and diabetes. Arteriosclerosis 10: 493-496. doi: 10.1161/01.ATO.10.4.493. PubMed: 2196039.

Hocking S, Samocha-Bonet D, Milner KL, Greenfield JR, Chisholm DJ (2012) Anthropometric correlates of total body fat, abdominal fat and visceral and subcutaneous adiposity in children and adolescents: National and Regional Differences. Nutr J 88: 1263-1271. doi: 10.1038/nj.2012.95. PubMed: 22710928.

30. McGill HC Jr., McManus CA (2003) Starting earlier to prevent heart disease. JAMA 289: 2320-2322. doi: 10.1001/jama.289.17.2320. PubMed: 14600192.

Cook S, Wiltzmann M, Auinger P, Nguyen M, Dietz WH (2004) Prevalence of a metabolic syndrome phenotype in adolescents: findings from the third National Health and Nutrition Examination Survey, 1988-1994. Arch Pediatr Adolesc Med 157: 821-827. doi: 10.1001/archpedi.157.8.821. PubMed: 12912790.

Johnson WD, Kroon JJ, Greenway FL, Bouchard C, Ryan D et al. (2009) Prevalence of risk factors for metabolic syndrome in adolescents: National and Regional Differences. Nutr J 8: 156. doi: 10.1186/1475-2891-8-156. PubMed: 19349567.

Steinberger J, Daniels SR, Eckel RH, Hayman L, Lustig RH et al. (2009) Progress and challenges in metabolic syndrome in children and adolescents: a scientific statement from the American Heart Association Atherosclerosis, Hypertension, and Obesity in the Young Committee of the Council on Cardiovascular Disease in the Young, Council on Cardiovascular Nursing, and Council on Nutrition, Physical Activity, and Metabolism. Circulation 119: 626-647. doi: 10.1161/CIRCULATIONAHA.110.191394. PubMed: 19139390.

Taksali SE, Caprio S, Dziura J, Dufour S, Call AM et al. (2008) High visceral and low abdominal subcutaneous fat stores in the obese adolescent: a determinant of an adverse metabolic phenotype. Diabetes 57: 367-371. PubMed: 17977954.
49. Siegel MJ, Hildebolt CF, Bae KT, Hong C, White NH (2007) Total and intraabdominal fat distribution in preadolescents and adolescents: measurement with MR imaging. Radiology 242: 846-856. doi:10.1148/radiol.2423060111. PubMed: 17244720.

50. Demerath EW, Shen W, Lee M, Choh AC, Czerwinski SA et al. (2007) Approximation of total visceral adipose tissue with a single magnetic resonance image. Am J Clin Nutr 85: 362-368. PubMed: 17284730.

51. Shen W, Chen J, Kwak S, Punyanitya M, Heymsfield SB (2011) Between-slice intervals in quantification of adipose tissue and muscle in children. Int J Pediatr Obes 6: 149-156. doi: 10.3109/17477166.2010.486833. PubMed: 20528126.

52. Demerath EW, Sun SS, Rogers N, Lee M, Reed D et al. (2007) Anatomical patterning of visceral adipose tissue: race, sex, and age variation. Obesity (Silver Spring) 15: 2984-2993. doi:10.1038/oby.2007.356. PubMed: 18198307.

53. Demerath EW, Reed D, Rogers N, Sun SS, Lee M et al. (2008) Visceral adiposity and its anatomical distribution as predictors of the metabolic syndrome and cardiometabolic risk factor levels. Am J Clin Nutr 88: 1263-1271. PubMed: 18996861.

54. Cornier MA, Marshall JA, Hill JO, Maahs DM, Eckel RH (2011) Prevention of overweight/obesity as a strategy to optimize cardiovascular health. Circulation 124: 840-850. doi:10.1161/CIRCULATIONAHA.110.968461. PubMed: 2184090.