Constitutional thinness and anorexia nervosa: a possible misdiagnosis?

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INTRODUCTION

In a young population of women, between 15 and 30 years old, a low body mass index (BMI) associated with apparent healthy state suggests that the diagnosis of anorexia nervosa (AN). For the same age range, other etiologies of starvation are associated with specific symptoms leading to obvious diagnosis of blood or oncologic pathologies, digestive absorption disorders.

Although restrictive anorexia nervosa (R-AN) definition reported in the “diagnostic and statistical manual of mental disorders” (DSM) is in perpetual evolution, diagnosis remains easy for an experienced clinician. The last issue, DSM5, was published in 2013 (1). Previous DSM IV definition (2) included some psychological elements in the diagnosis such as “refusal to maintain a normal BMI for their age and height, weight loss leading to major consequences of this truncated definition are further discussed as it can lead to misdiagnosis in front of a thin woman.

DIFFERENCES BETWEEN ANOREXIA NERVOSA AND CONSTITUTIONAL THINNESS

In a similar population of women, a low BMI associated with apparent healthy state could also suggests that the diagnosis of a not yet well-known entity called constitutional thinness (CT). This natural and physiological low BMI (<18.5 kg/m²) state is associated with preserved menses (without contraceptive treatment) and reproductive function leading to constitutional thin families. CT prevalence is unknown in the developed world. These subjects want to gain weight and often consult in this perspective.

Constitutional thinness and anorexia nervosa: a possible misdiagnosis. The results published by our group come from a large cohort of more than 600 R-AN and 100 CT.
Hormonal abnormalities data about R-AN reported so far in literature are commonly accepted and have been showed and published many times by different teams. Most of them are undernutrition markers such as low free-T3 (5,6), low IGF-1, and elevated GH displaying a GH resistance (7), elevated SHBG (8), and blunted leptin reflecting a decreased fat mass (9). These biological features are adaptive to food restriction and hormonal levels return within normal range after refeeding (10, 11). In 2010, we published biological data on a large R-AN cohort of 200 patients. Despite a correlation between BMI and five specific nutritional or hormonal parameters including leptin, GH, cortisol, free-T3, and IGF-1, we noticed a large inter-individual heterogeneity of these parameters (12). In CT matched for BMI and gender, free-T3, IGF-1, mean 24-h GH levels (sampled six times over the day) are in the normal range (Figure 1). This absence of undernutrition markers is in line with the low levels of leptin, still higher than in AN, and with preserved circadian cycle (13). A girl with low BMI and normal free-T3 or IGF-1 is not a R-AN.

Anorexia nervosa is associated with well-known hypothalamic–pituitary–peripheral axis changes/disturbances such as "pseudo-Cushing syndrome" and hypogonadism. Indeed, the excess of the hypothalamic–pituitary–adrenal (HPA) axis activity includes high CRF (14), increased circadian cortisol levels (15), and rapid escape of cortisol from suppression in response to i.v. dexamethasone (16). Nevertheless, R-AN patients display no clinical Cushing-like features and is considered as a “pseudo-Cushing syndrome.” These abnormalities regress after refeeding (17).

Restrictive anorexia nervosa amenorrhea, the only somatic trait in the DSM IV classification, is the consequence of blunted HPG axis associating low estradiol (18), low free testosterone levels (8), and absence of LH pulsatility (19). This low GnRH activity seems to be dependent on multiple neuropeptides/monoamines influences including opioid (20), gabaergic (21), or serotonergic activity (22) changes. Refeeding restores menses only in 55–100% of R-AN women (23–25). GnRH pump is the current validated treatment to restore LH pulsatility and to stimulate ovulation (26). This treatment leads to ovulation in 100% of cases, most often between day 10 and 14, and to pregnancy in 55% of cases after 6 stimulations (unpublished data).

Oppositely, normal HPA axis activity suggested by normal six-point (8, 12, 16, 20, 24, 04 h) circadian levels of cortisol was found in CT subjects matched for BMI and gender. Functional HPG axis was also attested by normal LH, FSH, estradiol, and free testosterone plasma level without oral contraception, and preserved fertility (13). A young adult girl with low BMI and normal menses without pills, and normal hormonal profile is not a R-AN.

The alteration of bone quality in AN, a consequence to undernutrition and hormonal abnormalities, is widely accepted. Loss of bone density, correlated with the duration of low BMI (27) and explained by a bone turnover uncoupling (28) leads to increased fractures risk (29), also related with some genetic disturbances (30). An important issue comes from the lack of treatment to increase bone mineral density, except refeeding. Hormonal deficiency supplementation with IGF-1 (31), classical treatment with bisphosphonates (32), or PTH administration (33) are not approved. The role of estradiol in osteoporosis treatment is worldwide accepted, but in R-AN the supplementation is non-effective (34). The association of two hormonal supplementations without side effects such as SDHA and estradiol seems to be effective (35, 36). However, all authors agree on the interest to treat earlier in order to preserve and not to restore bone mineral density.

Bone loss with 44% of osteopenia is also found in 20 years old CT subjects (37). Oppositely to the bone uncoupling seen in R-AN, bone turnover is balanced in CT, with normal bone formation (normal circadian osteocalcin profile) and bone resorption (normal plasma circadian CTX profile). Furthermore, bone microarchitecture measured by pQCT is different (37). Combined normal bone marker and specific microarchitecture profile are other arguments to differentiate CT from R-AN and to avoid a misdiagnosis.

While low food intake characterizes R-AN subjects, equilibrated energy balance was noticed in CT (38). Food restriction behavior of R-AN patients was found to be in opposition with the theoretical action of most of the appetite regulating hormones. Orexigenic gastric hormone ghrelin and the related gene derived peptide obestatin are elevated in all studies without orexigenic effect (13, 39). Ghrelin gene variants (40) or circulating antibodies (41) are proposed to explain the "ghrelin-resistance." Interestingly, a normal ghrelin level is found in lean anorectic patients (AN) with binging and in normal weight bulimic patients. High ghrelin seems to be a mark of restrictive subtype of AN (restrictive only and binging associated) (42). A recently discovered hypothalamic orexigenic peptide called 26 RFa is also increased in R-AN, and oppositely normal in CT (43). These hormones act through

| Anorexia nervosa | Constitutional thinness |
|------------------|-------------------------|
| BMI ≤ 17.5 kg/m² | No eating disorders; de-restriction |
| Psychological disorders | No psychological disorders |
| Amenorrhea | Physiological menses |
| Hormonal abnormalities= undernutrition | No hormonal abnormalities = no sign of undernutrition |
| T3 ↓, leptine ↓, IGF-1 ↓, GH ↑ | |
| Cortisol↑, ACTH ↑, 17 β E2 ↓, LH ↓, FSH ↓ | |
| Blunted fat mass | Diminished fat mass |
| Negative energy balance | Equilibrated / positive energy metabolism |
| Weight loss / broken weight growth curve | Stable weight within lower percentile of growth curve |

**FIGURE 1 | Psychiatric, hormonal, and energy balance differences between CT and AN**
the orexigenic neuron and peptide NPY. Conflicting results were reported on pre-prandial plasma NPY (39). A low-circulating NPY could explain the non-effacy of orexigenic hormones. Recent work showed role of NPY was more complex (44). These results need confirmation by circadian assessment of NPY levels. NPY data are not available in CT yet.

In CT, ghrelin and obestatin are in a normal range (13, 39). Therefore, normal ghrelin levels in a pure R-AN patient are questionable. In this case, misdiagnosis of CT or purging type AN should be discussed (42).

Studies on anorexic appetite regulating hormones as PYY and GLP-1 in AN present with conflicting results (11, 45, 46). PYY level in R-AN was found within normal range in our experience (11), but elevated in another study (47), perhaps due to the misdiagnosis of CT. Indeed, we reported in CT a high-PYY level throughout the day (08, 12, 16, 20, 24, 04 h) (11). CT’s leptin was found lower than in controls but higher than in R-AN with preserved circadian variation. Finally, low level of leptin, another anorexic peptide, found in all R-AN studies (13, 48) is probably more informative on the fat mass than on the eating behavior. Discrepancies found in appetite regulating hormones are in line with differences found in questionnaires measuring restrained eating behavior (DEBQ) or shape concern (EDE). Many questionnaires were proposed for eating disorders phenotyping. EDI questionnaire is a self-report questionnaire used to assess the presence of eating disorders, anorexia nervosa both restricting and binge-eating/purging type (49). EDE questionnaire deals with the frequency in which the patient engages in behaviors indicative of an eating disorder over a 28-day period (restraint, eating, shape, weight concern) (50), while anorexia nervosa both restricting and binge-eating/purging type (49). EDE questionnaire deals with the frequency in which the patient engages in behaviors indicative of an eating disorder over a 28-day period (restraint, eating, shape, weight concern) (50). DEBQ was developed to measure emotional external and restrain eating (51). All of these psychological scales present pathological scores in AN but not in CT. Moreover, we found an unrestrained eating behavior in CT when compared to controls (3).

For the same, low BMI, anorectic behavior, and adaptive appetite regulating hormone characterizes R-AN patients and distinguishes them from CT patients.

Finally, genetic pathophysiological role in AN was proposed throughout polymorphism gene studies (52) or familial histories of disease (53). Recently, two studies reported a tendency to the gene hypothesis without the statistical significantly in 421 probants for the first and 1606 probants for the second (54, 55). Significant ratio of chromosome 16p11.2 region duplications was previously noticed in lean patients with autism or schizophrenia (56), without any argument to relate these results to leanness rather than to psychiatric disorders. No genetic signature can be proposed at this moment for the CT patients despite the family pedigree (38). Some studies are in progress but no results are available. Along with all these data on genetic approach, it is important to underline the role of lean subjects’ phenotyping within these large cohorts’ studies.

**DISCUSSION**

This review focused on biological differences between R-AN and CT. As CT display specific clinical features and normal hormonal parameters, the misdiagnosis between CT and R-AN is no longer permitted. A very thin girl who claims a normal diet should be heard and not considered as a “liar” AN patient. A biological assessment including leptin, free-T3, and IGF-1 is proposed in order to avoid a misdiagnosis.

This review also focused on conflicting literature data reported in R-AN. Because of the clinical or biological heterogeneity of patients selected for the studies, publications results cannot always be compared and could explained the conflicting data. Therefore, straight and objective classification also called “phenotyping” is required in clinical research in order to obtain better comparable symptoms. For example, the definition shift between DSM IV and DSM5 changes the developing risk of AN, double in a dancer population (57) or in general population (58). Biological assessment including hormonal, nutritional markers, neuroimaging, or questionnaires could help on AN phenotyping. Currently, no biological determination or questionnaire evaluations are required for AN definition in DSM5.

In conclusion, CT represents a well-defined real state of low BMI associating a real weight gain desire, normal nutritional markers except for a mild decreased leptin, a constitutive appetite regulating hormone profile, the presence of menses in young women and low bone mass. While CT diagnosis is still poorly known, the new DSM5 AN definition proposing only psychological traits and no organic symptom is warring. In line with mentioned somatic differences, we advocate complementary biological markers in AN definition in order to avoid the misdiagnosis between AN and CT.

**ACKNOWLEDGMENTS**

The authors would like to thank Prof. Lang for his psychiatric perspective on this issue.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 03 July 2014; accepted: 03 October 2014; published online: 20 October 2014. Citation: Estour B, Galusca B and Germain N (2014) Constitutional thinness and anorexia nervosa: a possible misdiagnosis? Front. Endocrinol. 5:175. doi: 10.3389/fendo.2014.00175

This article was submitted to Neuroendocrine Science, a section of the journal Frontiers in Endocrinology.

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