Presentation, progression, and predictors of ovarian insufficiency in classic galactosemia

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Abstract
Classic galactosemia (CG) is an inherited metabolic disorder that affects about 1 in 50,000 live births in the United States and many other countries. With the benefit of early detection by newborn screening and rapid dietary restriction of galactose, generally achieved by removing dairy from the diet, most affected infants are spared the acute and potentially lethal symptoms of disease. Despite early detection and life-long dietary intervention, however, most patients grow to experience a constellation of long-term complications that include premature ovarian insufficiency in the vast majority of girls and young women. Our goal in the study reported here was to define the presentation, progression, and predictors of ovarian insufficiency in a cohort of 102 post-pubertal girls and women with CG. To our knowledge, this is the largest cohort studied to date. We found that 68% of the girls and women in our study achieved spontaneous menarche, while 32% achieved menarche only after starting hormone replacement therapy (HRT). Of those who achieved spontaneous menarche, fewer than 50% were still cycling regularly after 3 years, and fewer than 15% were still cycling regularly after 10 years. Of factors tested for possible association with spontaneous menarche, only detectable (≥ 0.04 ng/mL) plasma anti-Müllerian hormone (AMH) level was significant. These results extend substantially from prior studies and confirm that detectable plasma AMH is a useful predictor of ovarian function in girls and women with CG.

Introduction

Primary ovarian insufficiency (POI) is one of the most common long-term complications associated with classic galactosemia (CG), impacting a large majority of girls and women with CG, despite early detection (Fridovich-Keil et al. 2011; Berry 2014) and careful life-long dietary restriction of galactose (Frederick et al. 2017). Details of the presentation and progression of POI in CG have remained unclear, however, as has clarification of a useful prognostic marker for this outcome (Welling et al. 2017).

A literature trail extending back more than 35 years (Thakur et al. 2018) documents the striking prevalence of POI in CG. For example, Kaufman and colleagues (1981) reported that, of 15 girls and women aged 13–29 years in their study, 12 (80%) showed signs of POI. Waggoner et al. (1990) reported that 8 of 34 (or almost 24%) women in their study who were at least 17 years old experienced primary amenorrhea, and many more experienced later signs of POI. Schweitzer et al. (1993) reported that 5 of 11 (or almost 45%) girls and women in their study who were at least 17 years old experienced primary amenorrhea, and many more experienced later signs of POI. Spencer and colleagues (2013) reported that 11 of 28 (or 39%) post-pubertal women in their cohort had achieved menarche only after receiving hormone replacement therapy (HRT). Finally, van Erven and colleagues (2017) reported that > 55% of their cohort of 85 women with CG and current POI had required HRT to achieve menarche. In all of these studies, POI is a strikingly common experience for girls and women with CG. The apparent differences in prevalence reported may reflect the small sample sizes of most of the
studies, or differences in the study population, ascertainment, age cut-offs, and/or other differences.

The course of longitudinal progression of POI among girls and women with CG who achieve spontaneous menarche has also remained unclear. For example, of the 12 women with abnormal gonadal function in the study by Kaufman et al. (1981), fewer than half experienced primary amenorrhea, and most developed secondary amenorrhea or oligomenorrhea in their teens or 20s. Of the 26 women who achieved spontaneous menarche in the cohort of Waggoner et al. (1990), most developed oligomenorrhea or secondary amenorrhea within a few years, and only 5 of 17 were still cycling regularly in their early 20s. The studies by Schweitzer et al. (1993), Spencer et al. (2013), and van Erven et al. (2017) did not report on longitudinal progression of POI in their cohorts.

Finally, reliable predictors of POI in CG have remained unclear. For many decades, follicle stimulating hormone (FSH) was considered the marker of choice (e.g., Kaufman et al. 1981); however, studies have demonstrated that, in young girls who are not yet peri-pubertal, FSH can be unreliable (Sanders et al. 2009; Spencer et al. 2013). An alternative emerged when Sanders and colleagues (2009), and then Spencer and colleagues (2013), reported that plasma anti-Müllerian hormone (AMH), a glycoprotein produced by ovarian follicles at some but not all stages of development (La Marca et al. 2006; La Marca et al. 2010), was strikingly deficient in most girls and women with CG. AMH is now a well-established biomarker of “ovarian reserve” among girls and women who do not have galactosemia (Hagen et al. 2010; Hansen et al. 2011), but how accurately plasma AMH predicts ovarian function among girls and women with CG has remained uncertain. Specifically, Spencer and colleagues (2013) explored this question 5 years ago, but with relevant data from only 28 volunteers, they observed a compelling trend but lacked sufficient power to achieve a statistically significant result.

Here, we report analyses of the presentation, progression, and markers of ovarian function for 102 post-pubertal girls and women with CG. To our knowledge, this is the largest cohort reported to date. Our results extend substantially from prior studies and provide a foundation of knowledge supporting evidence-based predictions of ovarian function for girls and women with CG.

Materials and methods

Study volunteers

Cases for this study were selected from among volunteers already enrolled in our longitudinal protocol “Bases of Pathophysiology and Modifiers of Outcome in Galactosemia” (Emory IRB00024933; PI: JL Fridovich-Keil), which has been continuously approved by the Emory Institutional Review Board, or its predecessor, the Human Investigations Committee, since 1992. Volunteers for the larger study have been recruited via a combination of self-referral, predominantly from members of the Galactosemia Foundation (http://www.galactosemia.org), and referral from metabolic clinics, predominantly in North America. Post-pubertal girls and women with CG enrolled in the larger study were selected for inclusion here based on the availability of relevant marker and outcome information. Controls were post-pubertal unaffected siblings of cases in the larger study.

Outcomes quantified

Outcomes quantified for the current study included self-reported (for adults) or parent-reported (for children) information about age at menarche, whether menarche was spontaneous or HRT-assisted, and years of regular menstrual cycles following spontaneous menarche. We also collected information about HRT, where appropriate, including age at HRT initiation and termination, if relevant, and reasons for HRT initiation and termination, if relevant.

Markers and candidate covariates assessed

Markers tested for possible association with spontaneous menarche included AMH (in ng/mL) and FSH (in mIU/mL) quantified from plasma, as described previously (Sanders et al. 2009; Spencer et al. 2013). Unless otherwise noted, AMH and FSH values used in this study were derived from blood samples collected when the study volunteer was between 2 and 35 years old. Of note, while age does impact the normal range of AMH values (Lindhardt Johansen et al. 2013), the lower limit of that range for 2–35 year olds is well above the 0.04 ng/mL threshold of detection used here. Exogenous HRT shows only a small impact on AMH levels (La Marca et al. 2013), so AMH values used in this study were included whether or not the donor was on HRT at the time the blood was collected. In contrast, FSH level can be impacted dramatically by HRT, so the FSH values included here were restricted to those derived from samples collected when the donor was not receiving HRT. Where we had more than one relevant AMH or FSH value for a given volunteer, the appropriate values were averaged.

Covariates tested for potential association with spontaneous versus HRT-assisted menarche included race, year of birth, presence or absence of neonatal symptoms, family income, parent highest education level, and predicted residual GALT activity (Riehman et al. 2001).

Statistical analyses

Associations between spontaneous versus HRT-assisted menarche and potential covariates were tested through univariate
logistic regression models. Spontaneous menarche was considered the event of interest. Year of birth was considered as a continuous variable, whereas race, presence or absence of neonatal symptoms, family income, parent highest education level, and predicted residual GALT activity were considered as categorical variables. We used a likelihood ratio test to determine significance for all of the covariates; only year of birth was significant ($p < 0.05$). We then performed adjusted logistic regression models to test for association between spontaneous versus HRT-assisted menarche and AMH or FSH, with spontaneous menarche as the outcome of interest and continuous values of AMH or FSH as the independent variables, adjusting for year of birth.

To investigate years of continued spontaneous menstrual cycles beyond puberty, we created a Kaplan–Meier survival curve. The “event” quantified was cessation of spontaneous regular menstrual cycles. All calculations were conducted in SAS 9.4.

**Results**

**Prevalence of spontaneous versus HRT-assisted menarche among girls and women with CG**

Our cases for this study included 102 post-pubertal girls and women with CG and 22 controls, recruited as described in the Materials and methods section. The demographic characteristics of these volunteers are presented in Supplemental Table 1.

Of the cases in this study, 69 (68%) achieved spontaneous menarche and 33 (32%) achieved menarche only after starting HRT. The average age at spontaneous menarche in this cohort was 13.8 years (range 10 to 19), and the average age at HRT-assisted menarche was 14.2 years (range 11 to 18) (Fig. 1). As expected, 22 of 22 controls achieved spontaneous menarche at an average age of 12.8 years (range 9–17).

**Years of continued spontaneous menstrual cycles beyond puberty for young women with CG**

As a simple measure of continued ovarian function beyond puberty, we asked study volunteers who had achieved spontaneous menarche about years of continued regular cycling, without HRT, following puberty; we used these data to create a Kaplan–Meier “survival” curve. As illustrated in Fig. 2, only about half of those who achieved spontaneous menarche were still cycling regularly after 3 years, only about 35% were still cycling regularly after 5 years, and fewer than 15% were still cycling regularly after 10 years. Given that only 68% of the study volunteers had achieved spontaneous menarche to begin with, this means that, by their late teens, only about one-third of young women with CG in the study were cycling regularly without HRT, and by their mid-20s, the number was down to about 10%. Of note, the final sample size of women contributing longitudinal data for this calculation was 48.

**Plasma AMH but not FSH associates with spontaneous menarche in CG**

We tested both plasma AMH and FSH for a possible association with spontaneous menarche in our cohort. AMH was significant ($p = 0.0009$), as was AMH converted to a log$_{10}$ scale for sensitivity at the low end ($p = 0.0220$). Of note, from box and whisker plots of the data (Fig. 3a), it is clear that undetectable AMH was the most common result for volunteers from both the HRT-assisted and spontaneous menarche cohorts. An undetectable AMH result is, therefore, not particularly predictive. However, a detectable AMH level ($\geq 0.04$ ng/mL) appears to be highly predictive, as those scores were clearly skewed toward the spontaneous menarche group. Unlike AMH, FSH did not associate in a meaningful way ($p = 0.7871$) with spontaneous versus HRT-assisted menarche in our cohort (Fig. 3b).

**Starting and stopping HRT**

Of 55 girls and women with CG in the study who reported having taken HRT and who also gave a reason why HRT was initiated, 3 (5.4%) said they started HRT to stimulate growth and development, 27 (49.1%) said they started HRT to initiate or complete puberty, and 25 (45.5%) said they started HRT only after puberty to help manage irregular periods or other symptoms of menopause. The average age at initiation of HRT (Fig. 4) for the first group was 12.3 years (range 11–14), for the second group was 12.8 years (range 9–17), and for
the final group was 19.6 years (range 15–34). Nine additional women also reported taking HRT but for two of them, the reason for initiation was “other” and for seven, no reason was given.

Of the 64 women in the study who reported having ever taken HRT, 58 shared if they had stopped treatment, and, if so, at what age and why. Of these 58 women, 47 (81%) said they were still on HRT. Of 11 women (19%) who had stopped, the most common reason given was concern about possible health risks or side effects. Other reasons included “unknown”, “stopped working”, “hysterectomy”, and “elected not to use”. The average age of stopping HRT was 30.8 years (range 16–51). Of volunteers who were still taking HRT, the average age was 24.9 years (range 12.8–41.2).

Testing covariates for possible association with spontaneous menarche in CG

Finally, we tested each of six potential covariates for association with spontaneous menarche. These included the presence of acute neonatal symptoms before diagnosis, year of birth, race, and two indicators of family socioeconomic status (family income and parent’s highest level of education). Of these, only year of birth was even nominally significant (p = 0.0389), with girls born earlier being slightly more likely to achieve spontaneous menarche than girls born later. Specifically, for each year earlier a girl was born, she was 1.044 times as likely to have achieved spontaneous menarche.

The last potential covariate tested was residual GALT activity predicted from the GALT genotype (Riehman et al. 2001). Of 21 volunteers who reported HRT-assisted menarche and for whom we could predict GALT activity, only 2 (or 9.5%) had at least 0.4% predicted residual GALT activity. In contrast, of 51 volunteers who reported spontaneous
menarche, and for whom we could predict GALT activity, 12 (or 23.5%) had at least 0.4% predicted residual GALT activity. While this > 2-fold difference was notable, it was not statistically significant in our cohort ($p = 0.1497$).

**Discussion**

The work presented here is important for two reasons. First, it both confirms and extends the results of prior, smaller studies documenting the prevalence and progression of POI in CG. Second, and contrary to current recommendation (Welling et al. 2017), we found that detectable plasma AMH ($\geq 0.04$ ng/mL) was strongly positively associated with spontaneous menarche ($p = 0.0009$), meaning, in at least some cases, it could be used to inform prediction. Finally, by testing and excluding a handful of possible covariates, the results presented here help to rule out factors such as family socioeconomic status and neonatal acute symptoms as major contributors to POI in CG. In the search for factors that underlie outcomes, it is also useful to identify those that do not.

**Predicting spontaneous menarche**

One of the questions asked by families of girls with CG is: “What is the chance my daughter will be able to achieve spontaneous puberty?” From the results presented here, we can say that, with no further information, this chance is about 68%. If plasma AMH is undetectable ($< 0.04$ ng/mL), which is the most common result in CG, 68% is still our best estimate. When the AMH is detectable, however, we can do better. For example, when AMH is between 0.04 and 0.99 ng/mL ($n = 12$ in our cohort), the point estimate from a binomial exact test is 91.7% with a 95% confidence interval of 73.6% to 99.8%. If the AMH is 1 ng/mL or greater ($n = 6$ in our cohort), the point estimate is 100% with a 95% confidence interval of 60.7% to 100%.

Of course, that undetectable AMH was the most common result for girls and women in both the spontaneous and HRT-assisted menarche groups while detectable AMH was clearly skewed toward spontaneous menarche raises the question as to why this is the case. The most likely explanation may be that the threshold of detection of the AMH assay used on our samples was limiting. Specifically, we hypothesize that some of the samples scored here as “undetectable AMH” actually had very low but non-zero levels of AMH that were below the threshold of detection of the assay applied, and the trace ovarian function producing those very low levels was sufficient to enable spontaneous menarche, at least in some girls. As higher sensitivity assays for AMH are developed, this will become a testable hypothesis.

**Experience with HRT**

Our results presented here also give insight into the timing and reasons for HRT initiation and termination among girls and women with CG. For example, as presented in the Results section, a small number of girls in our study initiated HRT to promote growth, but the vast majority were split almost evenly between needing help beginning or completing puberty and needing relief from peri-menopausal symptoms. As expected, the average age of girls in the first two groups was lower (12.4 and 12.7 years, respectively) than of women in the third group (19.6 years). That said, in some cases, girls as young as 15 years started HRT to help control irregular periods, hot flashes, or other symptoms (Fig. 4).

The decision of if and when to stop HRT has been a point of particular uncertainty and concern for many women with CG. Current recommendations (Berry 2014; Welling et al. 2017) do not address this point, and, indeed, of the 65 women in our study cohort who had initiated HRT, the majority reported that treatment was still ongoing. The optimal duration of HRT for women with CG who have experienced POI remains unclear, but recommendations offered for other groups of young women with POI (e.g., women with Turner syndrome or fragile X pre-mutation carriers) may apply. In those cases, the recommendation (Sullivan et al. 2016) is to continue HRT at least until the normal age of menopause (e.g., around 50 years old) unless contraindicated, for example, by a personal or family history of breast cancer.
Limitations of our study

While informative, this study also had a number of serious limitations. For example, our cohort size of 102 post-pubertal girls and women may be large for a rare disorder like CG, but, nonetheless, it limited our power to draw conclusions. Our cohort was also overwhelmingly North American, white non-Hispanic, well educated, and self-selected for families who chose to participate in a research study. How these results would compare in other geographies and demographic groups is unknown. Finally, because most of our study volunteers had not attempted pregnancy, we could not make any meaningful statements about success rates in that regard. Specifically, of 15 women with CG in our cohort who reported attempting pregnancy, 14 had achieved spontaneous menarche and 9 became pregnant. Of these 9, all of whom had achieved spontaneous menarche, only 1 received medical intervention before becoming pregnant. That 9 of 15 women with CG in our cohort who attempted pregnancy did become pregnant is encouraging and consistent with results reported recently by van Erven and colleagues (2017); however, our sample was small and may not be representative of the larger population.

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Compliance with ethical standards

Conflict of interest A. Frederick, A. Zinsli, G. Carlock, K. Conneely, J. Fridovich-Keil all declare that they have no conflict of interest.

Animal rights (IACUC) This article does not contain any studies with animal subjects performed by the any of the authors.

References


Hagen CP, Aksgaarde L, Sørensen K et al (2010) Serum levels of anti-Müllerian hormone as a marker of ovarian function in 926 healthy females from birth to adulthood and in 172 Turner syndrome patients. J Clin Endocrinol Metab 95:5003–5010


Supplemental Table 1: Demographics of volunteers in this study

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¹Only racial/ethnic groups represented in this study population are listed.