Estrogens regulate glycosylation of IgG in women and men

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The immunologic potency of IgG is modulated by glycosylation, but mechanisms regulating this process are undefined. A role for sex hormones is suggested by differences in IgG glycans between women and men, most prominently with respect to galactose. We therefore assessed IgG galactosylation in 713 healthy adults from 2 cohorts as well as in 159 subjects from 4 randomized controlled studies of endocrine manipulation: postmenopausal women receiving conjugated estrogens, raloxifene, or placebo; premenopausal women deprived of gonadal hormones with leuprolide and treated with estradiol or placebo; men deprived of gonadal hormones with goserelin and given testosterone or placebo; and men deprived of gonadal hormones with goserelin and given testosterone or placebo together with anastrozole to block conversion of testosterone to estradiol. Menopause was associated with an increase in agalactosylated IgG glycans, particularly in the most abundant fucosylated nonbisected (G0F) glycoform. Conjugated estrogens and raloxifene reduced G0F glycans in postmenopausal women, while in premenopausal women leuprolide increased G0F glycans in a manner reversed by estradiol. Among men, goserelin increased G0F glycans, an effect blocked by testosterone through conversion to estradiol. These results establish estrogens as an in vivo modulator of IgG galactosylation in both women and men, defining a pathway by which sex modulates immunity.

Introduction

The capacity of IgG to mediate immune responses is determined in part by two glycans within the Fc region that modulate binding to Fc receptors and complement (1). Each glycan can assume over 30 different forms (Figure 1), a diversity that affords the opportunity to fine-tune humoral immunity. For example, in patients with HIV, a shift toward proinflammatory IgG glycans correlates with more effective antiviral defense, while in rheumatoid arthritis glycan changes may contribute to the ability of IgG to trigger pathogenic inflammation (2–6). Glycoforms lacking terminal galactose (termed G0, denoting zero galactoses, see Figure 1) are particularly proinflammatory because they confer an enhanced ability to fix complement and engage the activating IgG receptor FcγRIIIa while simultaneously blocking antiinflammatory mechanisms mediated through sialylated and/or bigalactosylated (G2) glycans (2, 4, 7, 8).

Despite the immunological importance of IgG glycosylation, its regulation is poorly understood. Epidemiological data suggest that endocrine factors may play a role. Population studies reveal an increase in G0 glycans in midlife in women but not in men (9–12). During pregnancy, a marked
decline in G0 fraction accompanies the second and third trimesters, which reverses within 3 months of
parturition (13, 14). Estradiol, the primary circulating form of estrogen, and to a lesser extent proges-
terone exhibit correlations between spot hormone levels and IgG glycans, particularly with respect to
differential galactosylation (12). Thus, estrogens and potentially other hormones are plausible candi-
date modulators of IgG glycosylation, though the observational nature of available data leaves a direct
etiologic connection uncertain.

Interestingly, estrogen production is not limited to women. While the male testis elaborates only small
amounts of estrogens directly, estradiol is routinely generated from testosterone through the action of aroma-
tase (also termed estrogen synthase) (15). In men, estradiol participates in skeletal maturation, body fat regu-
lation, and sexual function (16). Thus, an effect of estrogens on IgG glycosylation could be physiologi-
cally relevant in men as well as women. Whether testosterone directly alters IgG glycans has not been explored.

The consequences of a role for sex hormones in the determination of IgG glycosylation are potentially
substantial. Levels of these hormones not only differentiate men and women but also vary widely within
an individual across the life span. Further, related medical interventions are common, including postmeno-
pausal replacement therapy, oral contraception, testosterone supplementation, and aromatase inhibition.
We therefore sought to test the effect of estrogens and testosterone on human IgG galactosylation in vivo.

Results

G0 IgG increases in association with menopause. We used liquid chromatography to assess IgG glycans in 126
male and 119 female adult blood donors, employing monogalactosylated (G1) glycans as a normalizing
factor (17). G0/G1 glycans increased with age (Figure 2A). Dividing G0 glycoforms into their two major
subclasses, fucosylated nonbisected G0 (G0F) and fucosylated bisected G0 (G0FB), we found that the
magnitude of age-related change was most prominent in the more abundant G0F component (Figure
2B). G0F/G1 glycans increased with age in both women and men (β per year of age = 0.003 ± 0.002, P
= 0.038 vs. β = 0.008 ± 0.002, P < 0.0001). Below 50 years of age, G0F/G1 glycans were significantly
lower in women than men (P < 0.0001, adjusted for age); this difference was no longer evident in older individu-
als. Correspondingly, employing ≥50 years of age as an interaction term in linear regression, we found a
prominent increase in G0F/G1 glycans in women only, consistent with a potential role for menopause (β
= 0.16 ± 0.05, P = 0.004; men β = −0.07 ± 0.05, P = NS) (Figure 2C). Neither women nor men showed
an age group–dependent increment in G0FB/G1 (both β = 0.00 ± 0.03, P = NS) (Figure 2C).

We confirmed this observation in a second cohort of 468 adult donors recruited from the Orkney
Islands, Scotland, employing purified IgG to quantitate glycans without G1 normalization (18). Again,
below 50 years of age, women exhibited a lower proportion of G0F IgG glycans than men (P < 0.001,
corrected for age); this difference was no longer observed at ≥50 years. Whereas G0F increased gradu-
ally with age in both women and men (β = 0.22 ± 0.04, P < 0.0001 vs. β = 0.26 ± 0.04, P < 0.0001),
only women exhibited an increase in G0F glycans at age 50 (β = 4.46 ± 1.16 P = 0.0002; men β = −0.41 ± 1.30 P = NS) (Figure 2D and Supplemental Figure 1; supplemental material available online with
this article; https://doi.org/10.1172/jci.insight.89703DS1). This higher-resolution method identified
a parallel increase in G0FB glycans, again for women only, although with a much smaller magnitude
of shift (Supplemental Figure 1).
Taking advantage of clinical data within the Orkney cohort, we compared premenopausal women (n = 112) with women who reported cessation of menses and who had never received hormone supplementation (n = 77). This correction strengthened the G0F increment (β for menopause = 6.74 ± 1.30, P < 0.0001, compared with β per year of age = 0.16 ± 0.04, P < 0.0001) (Figure 2E). Interestingly, in premenopausal
women for whom information was available concerning oral contraceptive use, self-reported current utilization (\(n = 29\)) compared with no oral contraceptive (\(n = 81\)) was associated with a modest reduction in G0F glycans corrected for age (\(\beta = –1.88 ± 0.91, P = 0.04\)).

Estrogen agonists reduce G0F glycans in postmenopausal and premenopausal women. To evaluate the specific contribution of gonadal hormones, we obtained blood samples from 58 postmenopausal women randomized to placebo, conjugated estrogens (0.625 mg daily), or raloxifene (60 mg daily), regimens employed for alleviation of postmenopausal vasomotor symptoms and/or osteoporosis (19). Both conjugated estrogens and raloxifene reduced G0F/G1 glycans compared with placebo \((P < 0.05)\) (Figure 3A). We then assessed IgG glycans in 21 healthy premenopausal women treated on an investigational basis with the gonadotropin-releasing hormone (GnRH) analog leuprolide to lower gonadal steroids to postmenopausal levels and then randomized to placebo or transdermal estradiol (0.075 mg/d) \((0.075 mg/d)\) (20). G0F/G1 glycans increased in placebo recipients compared with estrogen recipients \((P = 0.03, Figure 3B), while the cohorts became indistinguishable once the GnRH agonist effect faded sufficiently for normal menses to resume (Figure 3B). These results confirm that estrogens drive IgG galactosylation and implicate estrogen deficiency in the increase in G0F glycans that accompanies menopause.

Estrogens derived from testosterone promote IgG galactosylation in men. Compared with premenopausal women, men exhibit lower levels of circulating estrogens, principally as estradiol derived from testosterone by aromatase (21). To evaluate whether sex steroids alter IgG glycans in men, we tested samples from healthy male volunteers treated with the GnRH agonist goserelin, as employed for testosterone deprivation therapy in prostate cancer, followed by placebo or 5 g of testosterone gel daily (16). In placebo recipients, serum testosterone and estradiol levels remained in the prepubertal range, while testosterone restored both hormones to the midportion of the adult reference range (Figure 4A). Intriguingly, G0F/G1 glycans rose in placebo recipients but remained stable in testosterone recipients (Figure 4A). To evaluate whether this effect reflected testosterone itself or rather testosterone-derived estradiol, we tested samples from men receiving the same interventions in the presence of the aromatase inhibitor anastrozole (1 mg daily), a treatment that resulted in levels of estradiol remaining equally low in testosterone and placebo groups (Figure 4B). Under these conditions, testosterone no longer normalized G0F/G1 glycans, confirming that its effect on IgG galactosylation was mediated not directly by testosterone itself but instead by testosterone-derived estradiol (Figure 4B).

Discussion

The relationship between sex and immune function is complex. Not all immunological differences between men and women are hormonal. For example, genes of immune relevance are found on the sex chromosomes (22). However, multiple immune lineages, including B cells, express estrogen and androgen receptors, implicating gonadal steroids in immune regulation (23). Understanding the role of hormones in immune function is important because these effects will give rise to immune differences between men and women as well as in association with childhood, puberty, pregnancy, and menopause. Further, an effect of sex hormones on immune function could lead to unintended consequences of the therapeutic manipulation of gonadal steroids.

In this study, we show that estrogens modulate human IgG galactosylation in vivo. Measurable changes in IgG glycans occur during menopause and from administration of hormones at therapeutic doses, including postmenopausal hormone replacement, oral contraception, and testosterone supple-
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This mechanism is operative in men as well, reflecting the action of estradiol derived from aromatization of testosterone rather than testosterone itself. These observations identify a pathway by which sex, as well as related medical interventions, affect immunity.

Altered IgG galactosylation is of particular interest because of the pivotal role of galactose on IgG effector function. G0 glycans enable IgG to engage mannose binding lectin, enhancing complement fixation via the lectin pathway (4, 24). G0 glycans also enhance the affinity of IgG for FcγRIIIa and therefore promote antibody-dependent cellular cytotoxicity, an effect implicated in antiviral potency (2). Whereas G1 glycans remain relatively stable across the adult age range, the increase in G0 glycans translates into a reduction in G2 glycans that engage the inhibitory IgG receptor FcγRIIb and dectin-1 to block chemoattractant signaling (8, 25). Moreover, only galactosylated Fc glycans are able to bear terminal sialic acid (see Figure 1), a modification that impairs Fc receptor binding while engaging a range of antiinflammatory mechanisms (7, 26–28). These considerations suggest that estrogens are likely to constrain the inflammatory capacity of the IgG pool.

One evolutionarily critical milieu in which the effect of estrogen on IgG could play a role is pregnancy. Pregnant women demonstrate a marked decrease in IgG G0 glycans (13, 14). Estradiol levels increase as much as 100-fold by the third trimester of pregnancy, such that the mechanism identified here is likely to be an important driver of the pregnancy-associated decline in agalactosylated IgG (29). Complement fixation is implicated in fetal demise, raising the possibility that estrogen-mediated glycan changes promote successful gestation (30).

Hormonal effects on IgG glycans could also affect immune-mediated disease. Autoantibody-containing IgG immune complexes and complement fixation are implicated in diseases such as rheumatoid arthritis (31). Rheumatoid arthritis is strongly associated with an increase in G0 IgG glycans, even preceding disease

![Figure 4. Estrogens regulate IgG glycosylation in men. (A) Serum testosterone and estradiol levels at baseline and 12 weeks in men treated with goserelin acetate and placebo or testosterone and changes in G0F/G1 glycans in response to these hormonal manipulations (n = 20/group). (B) Serum testosterone and estradiol levels at baseline and 16 weeks in men treated with goserelin acetate plus anastrozole together with placebo or testosterone and changes in G0F/G1 glycans in response to these hormonal manipulations (n = 20/group). **P < 0.01, ***P < 0.001, paired 2-tailed t test. The leftmost panel excludes from display (but not from analysis or error bars) an outlier in the 16-week placebo group (testosterone 1,858 ng/dl).](http://insight.jci.org)
onset (3, 32, 33). Further, the extent of glycan aberrancy correlates with disease severity (34). In pregnancy, reduction in G0 glycans correlates with improvement of rheumatoid arthritis disease activity, reversing in the postpartum period at a time of high risk for disease resurgence (13). Elevation in G0 glycans also occurs in vasculitis, juvenile idiopathic arthritis, systemic lupus erythematosus, and inflammatory bowel disease as well as during normal aging (3, 11, 32, 35–38). Recent studies have directly implicated cytokine-driven changes in IgG glycosylation (in particular sialylation) in the ability of murine IgG to induce experimental arthritis, corresponding to lower levels of sialylation and galactosylation in both total and autoantigen-specific IgG1 among asymptomatic human subjects who proceeded to develop rheumatoid arthritis within 12 months (5, 6). The effect of estrogens on this disease biology remains to be determined. Despite the female predominance of rheumatoid arthritis, the effect of estrogens observed here would ordinarily be expected to render antibodies less inflammatory. Consistent with this possibility, the incidence of rheumatoid arthritis among women rises in the immediate postmenopausal years, and premature menopause is associated with higher disease risk. Postmenopausal hormone supplementation appears to afford minimal, if any, protection perhaps because IgG glycosylation represents only one of many factors associated with disease risk (39–43).

We recognize important limitations to our study. The data herein do not address the mechanism by which estrogens alter IgG glycans. Although ligation of estrogen receptors on B cells is an appealing candidate, estrogen receptors are widely distributed, and it is possible that the effect of estrogens on glycosylation of IgG is fully or partially mediated by the effect of other lineages on B cell maturation and function. We have not determined whether the effect of estrogens on galactosylation extends to proteins beyond IgG. Indeed, a more widespread phenomenon is suggested by a cross-sectional study of plasma glycans that identified a general increase in G0 forms in women over the age of 50 years compared with younger women, potentially broadening the physiologic effect of estrogen-driven galactosylation (44).

A further limitation stems from our methodology. Most of our results were obtained using HPLC to characterize glycans from whole serum or plasma. In this context, relative specificity of G0 and G1 glycoforms for IgG arises because most N-linked glycans outside of the sterically restricted IgG Fc region are more highly developed (i.e., G2 and larger) (17). Even within the G0 subtraction, IgG depletion studies show that, while the large majority of G0F glycans originate with IgG, a substantial fraction of the minor G0FB component arises from other circulating proteins (data not shown). This effect likely explains why our HPLC studies found an estrogen-related difference only in G0F, while a statistically significant but much smaller menopause-associated shift in G0FB glycans could be detected in the Orkney analysis of purified IgG (Supplemental Figure 1). We therefore suspect that the effect of estrogens also extends to the lower-frequency (and even more proinflammatory) G0FB glycoform, although the utilization of HPLC for our experimental samples rendered us unable to test this possibility directly.

For the same reason, our methodology allowed us to address only IgG galactosylation, despite the importance of features such as fucosylation and sialylation to IgG function. Gender differences have been noted to extend to these features as well (9, 12). Galactosylation is required for attachment of sialic acid (see Figure 1), and galactosylation and sialylation frequently exhibit marked colinearity (2, 12). It is therefore possible that, by providing more galactose substrate, estrogens could indirectly promote greater IgG Fc sialylation. Consistent with this possibility, women but not men manifest a decrease in IgG Fc sialylation around the transition between the fifth and sixth decade of life (9). Further study is required to assess whether gonadal hormones affect sialylation, fucosylation, and other glycan features independently of galactosylation.

In summary, the abundance of G0 IgG glycans increases in women in association with menopause. Studies in healthy women undergoing endocrine manipulation indicate that this shift is mediated, at least in part, by the ability of estrogen to promote IgG galactosylation. Estradiol derived from aromatization of testosterone exerts a similar effect on IgG glycans in men. Taken together, these findings confirm that estrogens regulate human IgG glycosylation in vivo and thereby establish a mechanism through which sex modulates immunity.

Methods

Study participants. We assessed IgG glycosylation patterns in 6 cohorts. (a) The first cohort consisted of 245 blood donors from Brigham and Women’s Hospital, with samples obtained as discarded samples characterized only by age and gender. (b) The second cohort consisted of 468 consented adult volunteer participants in the Orkney Complex Disease Study (ORCADES), Orkney Islands (18, 32). (c) The third cohort consisted of 58 healthy postmenopausal women (age 56 ± 4 years, age at menopause 48 ± 6 years) randomized to oral therapy with placebo, conjugated estrogens (Premarin, Pfizer Inc.) (0.625 mg daily), or raloxifene...
(Evista, Eli Lily Inc.) (60 mg daily) for 18 months, selected at random from a larger study (n = 119) of the metabolic effects of hormone replacement (19). Blood samples were collected at baseline and at the end of the intervention. Women randomized to conjugated estrogens whose uteri were intact also received medroxyprogesterone acetate (5 mg daily) for 13 consecutive days trimonthly for endometrial protection; placebo and raloxifene subjects received cyclic placebo. (d) The fourth cohort consisted of 21 healthy premenopausal women (age 34 ± 7 years) treated with 5 monthly intramuscular doses of the GnRH analog leuprolide acetate (TAP Pharmaceutical Products Inc.) (3.75 mg) to suppress endogenous gonadal steroids and then randomized to concurrent treatment with transdermal estradiol (Bayer HealthCare Pharmaceuticals Inc.) (0.075 mg/d) or placebo. Patients were selected at random from a larger study (n = 70) of the metabolic effects of estradiol in premenopausal women (20). (e) The fifth cohort consisted of 40 healthy male volunteers (age 33 ± 7 years) treated with the GnRH agonist goserelin acetate (Zoladex, AstraZeneca Inc.) (3.6 mg subcutaneously at weeks 0, 4, 8, and 12) to suppress endogenous gonadal steroids and then randomized to simultaneous administration of placebo or testosterone gel (AndroGel, AbbVie Inc.) 5 g daily. Samples were analyzed for glycans at baseline and at 12 weeks. (f) The last cohort consisted of 40 healthy male volunteers (age 34 ± 6 years) treated with goserelin acetate (Zoladex, AstraZeneca Inc.) (3.6 mg subcutaneously at weeks 0, 4, 8, and 12) plus anastrozole (Arimidex, AstraZeneca Inc.) (1 mg orally daily) to block aromatization of testosterone to estradiol and then randomized to simultaneous administration of placebo or testosterone gel (5 g daily). Samples were analyzed for glycans at baseline and at 16 weeks. Cohorts e and f were selected at random from subjects in a larger study in which men were randomized to a range of doses of testosterone gel (placebo, 1.25, 2.5, 5, or 10 g daily), with (n = 198) or without (n = 202) administration of anastrozole; the 5 g testosterone group was selected as the comparator for this post-hoc study, because this dose maintained levels of gonadal steroids similar to baseline (16). Participants in all studies were unaware of study group assignment. Descriptions of the various cohorts, the interventions, and study results have been published (16, 19, 20, 45).

IgG glycan analysis. Samples from blood donors and hormonally manipulated subjects were analyzed using an established method (17, 32). Glycans were cleaved enzymatically from 5 μl of serum or plasma, conjugated to the fluorochrome 2-aminobenzamide, and quantitated using a TSKgel Amide-80 5-μm column attached to a Waters 1525 binary HPLC pump with a Waters 2487 dual wavelength detector, a Waters 2475 multijet fluorescence detector, and a Waters 717plus autosampler. To permit comparison across individuals, G0 glycans were normalized to the G1 fraction, which is stable across the adult age range (17, 25). In this method, selectivity for IgG glycans arises because the large majority of G0 and G1 forms in serum or plasma are from the IgG Fc region (17). Samples from ORCADES donors were analyzed from IgG purified by plate-bound protein G, using ultra performance liquid chromatography (UPLC), as described previously (18). For both methods, analysis was restricted to neutral glycans in order to avoid contamination from glycans attached to the Fab region of some IgG, since these are predominantly sialylated. HPLC studies were performed at the Brigham and Women’s Hospital, and UPLC studies were performed at the Genos Glycoscience Research Laboratory; within each comparison, samples were collected and analyzed simultaneously to avoid batch effects. The specific glycans fractions quantitated under HPLC and UPLC protocols are depicted in Supplemental Figure 2 and Supplemental Figure 3, respectively.

Statistics. Variance in glycoform abundance with age was assessed using Pearson statistics. Comparison of galactosylation in men and women, adjusted for age, employed general linear regression. An interaction term for age group (<50 vs. ≥50 years), menopause (yes/no by self-report), or oral contraceptive use (by self-report) was incorporated to assess associated increments in galactosylation (β coefficient ± SEM). Comparison among experimentally manipulated subjects was performed using Student’s t tests (2 tailed) for data normally distributed per Shapiro-Wilks and by Mann-Whitney U tests for nonparametric data. Error bars represent SEM. Statistical tests were performed using SAS 9.4 or GraphPad Prism 6 software. Significance was established at α = 0.05.

Study approval. The analyses reported here represent secondary use of previously collected samples, conducted under Brigham and Women’s Hospital institutional review board approval P2004000060.

Author contributions
AE and PAN conceptualized and designed the research. AE, PMR, and GL acquired data. AE, JC, and PAN analyzed data. WK, KDD, MP, EWY, JSH, HC, UBK, JFW, and JSF provided samples, reagents, and other project assistance. PAN wrote the manuscript, and all authors edited the manuscript.
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41. Salliot C, Bombardier C, Saraux A, Combe B, Dougados M. Hormonal replacement therapy may reduce the risk for RA in women with early arthritis who carry HLA-DRB1 *01 and/or *04 alleles by protecting against the production of anti-CCP: results from the ESPOIR cohort. *Ann Rheum Dis.* 2010;69(9):1683–1686.
