Book Chapter

Astrocytic Estrogen Receptors and Impaired Neurotrophic Responses in a Rat Model of Perimenopause

Todd E Morgan and Caleb E Finch*

Davis School of Gerontology, Department of Biological Sciences, Dornsife College, University of Southern California, USA

*Corresponding Author: Caleb E Finch, Davis School of Gerontology, Department of Biological Sciences, Dornsife College, University of Southern California, Los Angeles, CA, USA

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Abstract

In a perimenopausal model of middle-aged rats, the astrocyte estrogen receptor-alpha (ERα): ER-beta (ERβ) ratio increased with the onset of acyclicity (constant estrus, CE) in association with impaired neurotrophic responses to estradiol (E2). We report additional data on irregular cycling (IR) from this study of 9 month old perimenopausal subgroups. In particular, irregular cyclers (IR) also show increased ERα:ERβ ratio in cerebral cortex astrocytes comparable to acyclic individuals in CE. In mixed glial cultures from these same cycling subgroups, the E2-dependent neurotrophic activity and glial fibrillary acidic protein (GFAP) repression by E2 were impaired in IR to the same degree as in CE-derived glia. The greater importance of cycling status than age during the perimenopause to astrocyte ERs are attributable to individual variations of the residual ovarian follicle pool, which determine the onset of acyclicity. The corresponding loss of E2-dependent GFAP repression and E2-dependent neurotrophic activity add further to the inverse relationship of GFAP expression and astrocyte neurotrophic activity across aging in both sexes. These findings are relevant to impairments of spatial learning and of hippocampal long-term potentiation during the onset of IR in middle-aged rats, and to perimenopausal factors mediating the higher risk of women for Alzheimer disease.

Keywords

Estrogen Receptors; Perimenopause; Astrocytes; Glial Fibrillary Acid Protein; Neurotrophic
Introduction

Estrogen receptor-alpha (ERα) and estrogen receptor-beta (ERβ) undergo age-related shifts in both sexes [1-6]. In brain, ER aging changes vary by cell type and hormonal status. In a perimenopausal model of middle-aged rats, the astrocyte ERα:ERβ ratio was increased in acyclic constant estrus (CE) rats [4]. The ERα:ERβ ratio has broad significance to synaptic plasticity [6,7], which we have characterized for astrocytic neurotrophic support [3,4,8,9]. In particular, elevations of ERα:ERβ decrease neurotrophic support, which is indirectly linked to expression of glial fibrillary acidic protein (GFAP), the astrocytic intermediate filament through its estrogen response elements [9,10]. The elevated ERα:ERβ in astrocyte cultures from aging males was manipulated by RNAi, which decreased GFAP [4,8]. Both E2 responses (GFAP repression and induction of neurotrophic activity) are lost during aging [4,8]. In this review, we present data on irregular cycling (IR; Arimoto et al., [3]) that extends findings of Arimoto et al. [4] and discuss how changes in astrocytic ERs during the perimenopause transition may result from changes in steroids from both the ovary and the brain. The importance of IR status was shown in impaired learning of IR rats vs. RC (regular cycling) of the same age [11] and impaired long-term potentiation in hippocampal slices [12].

A group of 9 month old rats were characterized for cycling status: regular cycles 4–5 day (RC); irregular cycles >5 day (IR); and acyclic CE. The IR group was not reported in Arimoto et al. [4] but was described in Arimoto et al. [3] and the Ph.D. Thesis of Jason Arimoto (USC Department of Biological Sciences, 2012). First we describe astrocyte ER in vivo expression in 9 month old rats. Then we describe primary glial cultures from the same groups for astrocyte GFAP expression in relation to neurotrophic activity.

Astrocyte ERα:ERβ Ratio Increases with Age

Age changes in astrocyte estrogen receptors (ER) are under-represented in the growing literature on ERs in brain aging, which has mainly focused on neuronal ERs [7,13]. Both sexes of
aging rats (9–24 month, middle-age to old age) show increased ERα:ERβ in astrocytes of cerebral cortex as an outcome of increased ERα and decreased ERβ, as determined from immunohistochemical analysis of co-labeling for GFAP, the astrocyte-specific protein [4]. Figure 1 shows progressive increase of ERα and decrease of ERβ during transitions from RC to IR to CE (Figures 1A,B). In the prior comparison of CE with RC [4], ERβ levels were lower by 40%, whereas IR were just 10% lower (Figure 1B). Correspondingly, the ERα:ERβ ratio increased by 1.2-fold in IR and 2.5-fold in CE (Figure 1C). In an older cohort at 13 month, CE and IR had similar shifts of ERα:ERβ ([4], not shown). Primary glial cultures of enriched astrocytes or mixed glial (astrocytes: microglia, 3:1) retain these age changes. Perimenopausal rats age 9 month show further complexity in comparisons of regular cyclers (RC) vs. irregular cyclers (IR) vs. acyclic CE. The data on IR [3,5] show ERα:ERβ shifts in vivo that are comparable to the CE in 9 month old rats (Figure 1A). We suggest that astrocytes may be major contributors to the decrease of ERβ in whole hippocampal RNA from IR vs. RC rats [12].

**Figure 1:** (A–C) *In vivo* ERα and ERβ expression in cerebral cortex astrocytes of perimenopausal female rats, aged 9 month differ by ovarian cycle status: regular cycles (RC) 4–5 day irregular cycles (IR) >5 day; constant estrus (CE) [14,15]. Astrocyte ERs in primary visual cortex in layers 1 and 2/3, RC vs. IR vs. CE; 4 brains per cycle status; Data is expressed as % regularly cycling (RC) control; mean+SEM; ***p < 0.001; **p < 0.01; *p < 0.05. RC and CE data published in Arimoto et al. [4], Figure 7 (n.b. CE is denoted as AC, ibid.). The IR data are from Arimoto et al. [3] and the thesis of Jason Arimoto (USC, 2012). (D,E) *In vitro* GFAP expression and neurotrophic activity (MAP5,
neurite outgrowth) in mixed glia from 9 month old perimenopausal rats of different cycling status: RC vs. IR vs. CE. Mixed glial were co-cultured with E18 neurons; 4 cultures, 4 brains per cycling stage group. Data expressed as % RC control; mean + SEM; **p < 0.01; *p < 0.05. GFAP responses to 100 pM E2 in mixed glia from 9 month rats: The RC responded to E2 with decreased GFAP, but age-matched IR and CE were unresponsive. Neurite outgrowth (MAP5) was increased by 100 pM E2 in RC, but not in age-matched IR and CE. The RC and CE data were published in Arimoto et al. [4], Figure 6 (n.b. CE is denoted as AC, ibid.). The IR data are from Arimoto et al. [3] and the thesis of Jason Arimoto (USC, 2012). (F) Reciprocal relationships between GFAP and neurite outgrowth (MAP5), plotted as % change relative to controls in that experiment. RC and CE data published in Arimoto et al. [4], Figure 8. New data on 9 month IR are calculated from (D,E).

**Inverse Relationships between GFAP and Neurotrophic Activity**

Primary cultures of mixed glia (astrocytes:microglia, 3:1) were prepared from the other hemicortex from these rats [4]. Mixed glia were used rather than enriched astrocytes because we wished to avoid artifactual perturbation of gene activity associated with hydrodynamic effects of shaking to remove microglia [16]. In male derived glia, mixed glia and enriched astrocytes showed the same age trends in ERs and GFAP, and neurotrophic activity [4]. In glia from 9 month rats, astrocyte GFAP was repressed by E2 in RC, whereas IR and CE were unresponsive to E2 (Figure 1D). Correspondingly, neurotrophic activity in conditioned media assayed by neurite outgrowth was increased by E2 only in glia from regular cyclers (Figure 1E). The reciprocal relationship between GFAP expression and astrocyte neurotrophic activity (RC vs. IR vs. CE) is consistent with prior analysis, updated in Figure 1F.

Enhanced GFAP expression influences neurite outgrowth through astrocyte-secreted laminin and other extracellular factors [9]. Loss of E2-sensitivity of neurite outgrowth in male derived astrocytes was reversed by down regulation of GFAP by RNAi; conversely, young astrocytes acquired an aging phenotype by transfection with GFAP cDNA [8]. Because GFAP is regulated by E2 through a classical promoter ERE that binds ERα [10], we examined the role of ERs. The elevated ERα:ERβ in astrocyte cultures from aging males was manipulated by RNAi to ERα,
which decreased GFAP [4]. However, neurotrophic activity was not restored, suggesting additional ER dependent mechanisms. The loss of E2 responses in GFAP repression and in E2-dependent neurotrophic activity may be the earliest impairment in gene responses in a rodent model.

**Perimenopausal Transition**

The rodent perimenopausal model (Figures 2A,B) is described with reference to the STRAW stages of human menopause [15]. Rodents and humans undergo similar ovarian senescence, beginning with increasingly IR and declining fertility, and ending with total depletion of ovarian follicles [17]. The CE following IR with modest sustained blood E2 and very low progesterone (P4) may be considered as a model for hyperestrogenic cycles during human peri-menopause [15]). We hypothesize that the perimenopause may be characterized by a general trend for increasing plasma E2:P4 during the perimenopause [15,17], equivalent to progressive exposure to unopposed estrogen.

![Figure 2](image-url): (A) Comparison of perimenopausal transitions in women and rodents (lab mouse and rat). The most common initial acyclic state of rodents, CE is driven by continued ovarian estradiol (E2) production at low levels with...
hypothalamic impairments of the E2-dependent preovulatory gonadotrophin surge. Alternate pathways with persistent diestrus (PDE) are shown in panel B. The terminal state of anestrus with total oocyte depletion arise after 20 month, with LH elevations as in postmenopausal women. Rodent perimenopause occupies a larger fraction of the 28 month life span than for women, spanning the onset of cycle irregularity after 6 month to ovarian follicle depletion after 20 month, which comprises half the lifespan. For details, see [14,15,17-19]. (B) Rodent perimenopausal transitions with alternate trajectories from lengthening cycles interspersed with 4–5 day ovulatory cycles (I) CE followed by PDF (repetitive pseudopregnancy), and thence to anestrus; (II) CE directly to anestrus; (III) Lengthening cycles leading directly to anestrus (least common). Adapted from [17]. (C) Studies on C57BL/6J mice from the Finch lab. Ovarian oocyte numbers in Y (young, 4–5 month) and M (middle-aged, 13–14 month); the M were grouped by the occurrence of recent ovulation. Horizontal lines are median values. From Gosden et al. [20]. (D) Heterochronic ovary transplantation between Y and M ages. The MLovx were ovariectomized (OVX) at 3 month and given ovaries at 12 month; their aging without exposure to ovarian steroids maintained the potential for 4 day cycling, not seen in Y-M transplant. Redrawn from Felicio et al. [21].

To study perimenopausal changes associated with cycle lengthening, we used a mixed glia model (3:1 astrocytes; microglia), which had shown the same age changes of ERa, GFAP, and neurotrophic activity as enriched astrocytes in males [4]. Mixed glia from irregularly cycling 9 month rats and CE rats both show impaired E2-responses of GFAP and of neurotrophic activity. Thus, the loss of hypothalamic GFAP responses to E2 in middle-aged rats [22] has a counterpart in cerebral cortex astrocytes. Moreover, it shows the impact of modest steroidal perturbations associated with irregular cycles to synaptic functions related to memory. In IR vs. RC of the same age, hippocampal slices show impaired LTP [12], while intact rats show impaired spatial learning [11]. In an astrocyte-neuron culture model, ERa was specifically associated with E2-induced glutamatergic synaptogenesis [23]. We suggest that the impairments of E2-dependent neurotrophic activity in IR rats can be used to identify E2-dependent neurotrophic factors underlying LTP and spatial memory.

**Ovarian Contributions**

We suggest that the early onset of ER changes in females is driven by ovarian steroidal changes. In particular, as estrous cycles undergo lengthening and transition to acyclicity, there is a
modest reduction of the blood estradiol: progesterone ratio (E2:P4) [17]. At much later ages, typically after 20 month in rodents, ovarian follicles are depleted with reductions of plasma E2 to ovariectomized (OVX) levels [19]. Although male rodents do not undergo total reproductive senescence with the complete loss of fertility and gonadal steroids in older age, plasma testosterone tends to decrease in middle-aged male rats [24].

Further perspectives come from studies of 3 decades ago in our lab. By 12–14 month, rodent ovaries have < 10% of oocytes and primary follicles remaining from the young adult (Figure 2C; Gosden et al., [20]). Note the wide range of oocyte numbers; this 5-fold wide range is hypothesized to underlie the individual differences in onset of acyclicity [25].

The role of oocyte depletion in cycle lengthening was shown with ovarian transplants between young and middle-aged mice (heterochronic transplants; Figure 2D; Felicio et al., [21]). Replacing young ovaries with middle-aged ovaries caused premature cycles lengthening to >5 days. Despite the wide range of remaining follicles in middle-aged ovaries (Figure 2C), the remaining number is close to the threshold required for ovulatory cycles. The tight linkage of cycle lengthening to the remaining ovarian follicles is also shown by the induction of premature cycle lengthening and CE by surgically removing 90% of young ovarian mass [26].

**Neuroendocrine Contributions**

Neuroendocrine mechanisms are also at work, as shown by transplant of young ovaries to middle-aged mice, which yielded mostly 5 day cycles. However, if middle-aged mice were OVX when young and allowed to age without the presence of ovaries (long-term ovariectomy, Lt-OVX), then 4 day cycles were observed. Thus, the loss of 4 day cycles has a neuroendocrine component that is modified by estrogen exposure. The estrogen exposure hypothesis of cycle lengthening was tested by several durations of exposure to exogenous E2, which advanced the onset of acyclicity, in which the neuroendocrine component was shown by transplantation of control ovaries [27,28]. As few as 6
weeks of low sustained E2 accelerated the onset of acyclicity [28]. The importance of the E2:P4 ratio was shown by the protective effect of P4 implants [29]. We attribute these effects of E2:P4 on synaptic remodeling rather than neurodegeneration because chronic E2 did not change the number of hypothalamic LHRH or TIDA neurons [30]. Astrocyte GFAP is a mediator of estrous cycles by the close relation of GFAP-containing astrocyte processes to the LHRH neurons which shift during proestrus [31]. The E2-dependent induction of hypothalamic GFAP was impaired in middle-aged rats in parallel with the loss of the E2-induced LH surge [22]. Subsequently, we developed *in vitro* systems for studying GFAP regulation that further documents impaired GFAP responsiveness with functional connections to neuronal plasticity.

**Conclusion**

Laboratory rodents show a perimenopausal transition with lengthening cycles that has both brain (neuroendocrine) and ovarian contributions, as shown by prior ovarian transplantation studies between different age groups. Recent data further show that the neurotrophic activity of astrocytes assayed *in vitro* declines in the earliest perimenopausal stage of cycle irregularity and with cycle lengthening. These findings are relevant to impairments of spatial learning and of hippocampal long-term potentiation during the onset of IR and to perimenopausal factors mediating the higher risk of women for Alzheimer disease.

**References**

1. Wilson ME, Rosewell KL, Kashon ML, Shughrue PJ, Merchenthaler I, et al. Age differentially influences estrogen receptor (ERa) and estrogen receptor- (ERb) gene expression in specific regions of the rat brain. Mech. Ageing Dev. 2002; 123: 593–601.
2. Wu D, Lin G, Gore AC. Age-related changes in hypothalamic androgen receptor and estrogen receptor alpha in male rats. J. Comp. Neurol. 2009; 512: 688–701.
3. Arimoto JM, Lin SW, Iwata N, Morgan TE, Finch CE. Aging and stage of reproductive senescence affects E2
mediated neurite outgrowth. Program No. 596.11. in Neuroscience Meeting Planner, Washington: Society for Neuroscience. 2011. Available online at: http://www.abstractsonline.com/Plan/ViewAbstract.aspx?Key=d5e7469c-f520-49b3-a8cb-3486d3ad3039&cKey=7e806c52-5ca6-47ad-a23a-f80335ad3801&mKey=8334be29-8911-4991-8c31-32b32dd5e6c8

4. Arimoto JM, Wong A, Rozovsky I, Lin SW, Morgan TE, et al. Age increase of estrogen receptor alpha (ERα) in cortical astrocytes impairs neurotrophic support in male and female rats. Endocrinology. 2013; 154: 2101–2113.

5. Arimoto JM. Glial support of neurite outgrowth: effect of aging and estrogen. PhD Thesis (Dept. Molecular Biology, Dornsife College, University of Southern California, Los Angeles). 2012. Available online at: http://digitallibrary.usc.edu/cdm/ref/collection/p15799coll3/id/36906

6. Foster TC. Role of estrogen receptor and expression and signaling on cognitive function during aging. Hippocampus. 2012; 22: 656–669.

7. Bean LA, Ianov L, Foster TC. Estrogen receptors, the hippocampus and memory. Neuroscientist. 2014; 20: 534–545.

8. Rozovsky I, Wei M, Morgan TE, Finch CE. Reversible age impairments in neurite outgrowth by manipulations of astrocytic GFAP. Neurobiol. Aging. 2005; 26: 705–715.

9. Rozovsky I, Wei M, Stone DJ, Zanjani H, Anderson CP, et al. Estradiol (E2) enhances neurite outgrowth by repressing glial fibrillary acidic protein expression and reorganizing laminin. Endocrinology. 2002; 143: 636–646.

10. Stone DJ, Song Y, Anderson CP, Krohn KK, Finch CE, et al. Bidirectional transcription regulation of glial fibrillary acidic protein by estradiol in vivo and in vitro. Endocrinology. 1988; 139: 3202–3209.

11. Paris JJ, Walf AA, Frye CA. Cognitive performance of middle-aged female rats is influenced by capacity to metabolize progesterone in the prefrontal cortex and hippocampus. Brain Res. 2011; 1379: 149–163.

www.videleaf.com
12. Yin F, Yao J, Sancheti H, Feng T, Melcangi RC, et al. The perimenopausal aging transition in the female rat brain: decline in bioenergetics systems and synaptic plasticity. Neurobiol. Aging. 2015; 36: 2282–2295.
13. Kermath BA, Riha PD, Woller MJ, Wolfe A, Gore AC. Hypothalamic molecular changes underlying natural reproductive senescence in the female rat. Endocrinology. 2014; 155: 3597–3609.
14. Nelson JF, Felicio LS, Randall PK, Sims C, Finch CE. A longitudinal study of estrous cyclicity in aging C57BL/6J mice: I. Cycle frequency, length and vaginal cytology. Biol. Reprod. 1982; 27: 327–339.
15. Finch CE. The menopause and aging: a comparative perspective. J. Steroid Biochem. Mol. Biol. 2014; 142: 132–141.
16. Gatson JW, Simpkins JW, Yi KD, Idris AH, Minei JP, et al. Aromatase is increased in astrocytes in the presence of elevated pressure. Endocrinology. 2011; 152: 207–213.
17. Finch CE, Felicio LS, Mobbs CV, Nelson JF. Ovarian and steroidal influences on neuroendocrine aging processes in female rodents. Endocr. Rev. 1984; 5: 467–497.
18. Felicio LS, Nelson JF, Finch CE. Longitudinal studies of estrous cyclicity in aging C57BL/6J mice: II. Cessation of cyclicity and the duration of persistent vaginal cornification. Biol. Reprod. 1984; 31: 446–453.
19. Gee DM, Flurkey K, Finch CE. Aging and the regulation of luteinizing hormone in C57BL/6J mice: impaired elevations after ovariectomy and spontaneous elevations at advanced ages. Biol. Reprod. 1983; 28: 598–607.
20. Gosden RG, Laing SC, Felicio LS, Nelson JF, Finch CE. Imminent oocyte exhaustion and reduced follicular recruitment mark the transition to acyclicity in aging C57BL/6J mice. Biol. Reprod. 1983; 28: 255–260.
21. Felicio LS, Nelson JF, Finch CE. Prolongation and cessation of estrous cycles in aging C57BL/6J mice are differentially regulated events. Biol. Reprod. 1986; 34: 849–858.
22. Anderson CP, Rozovsky I, Stone DJ, Song Y, Lopez LM, et al. Aging and increased hypothalamic glial fibrillary acid protein (GFAP) mRNA in F344 female rats. Dissociation of
GFAP inducibility from the luteinizing hormone surge. Neuroendocrinology. 2002; 76: 121–130.

23. Jelks KB, Wylie R, Floyd CL, McAllister AK, Wise P. Estradiol targets synaptic proteins to induce glutamatergic synapse formation in cultured hippocampal neurons: critical role of estrogen receptor-alpha. J. Neurosci. 2007; 27: 6903–6913.

24. Rosario ER, Chang L, Beckett TL, Carroll JC, Paul Murphy M, et al. Age-related changes in serum and brain levels of androgens in male Brown Norway rats. Neuroreport. 2009; 20: 1534–1537.

25. Finch CE, Kirkwood TB. Chance, Development and Aging. New York: Oxford press. 2000.

26. Nelson JF, Felicio LS. Radical ovarian resection advances the onset of persistent vaginal cornification but only transiently disrupts hypothalamic-pituitary regulation of cyclicity in C57BL/6J mice. Biol. Reprod. 1986; 35: 957–964.

27. Mobbs CV, Kannegieter LS, Finch CE. Delayed anovulatory syndrome induced by estradiol in female C57BL/6J mice: age-like neuroendocrine, but not ovarian, impairments. Biol. Reprod. 1985; 32: 1010–1017.

28. Kohama SG, Anderson CP, Osterburg HH, May PC, Finch CE. Oral administration of estradiol to young C57BL/6J mice induces age-like neuroendocrine dysfunctions in the regulation of estrous cycles. Biol. Reprod. 1989; 41: 227–232.

29. Kohama SG, Anderson CP, Finch CE. Progesterone implants extend the capacity for 4-day estrous cycles in aging C57BL/6J mice and protect against acyclicity induced by estradiol. Biol. Reprod. 1989; 41: 233–244.

30. Kohama SG, Brown SA, Finch CE, McNeill TH. Chronic estradiol administration did not cause loss of hypothalamic LHRH or TIDA neurons in young or middle-aged C57BL/6J mice. Brain Res. 1992; 574: 341–344.

31. Garcia-Segura L, Luquin S, Párducz A, Naftolin F. Gonadal hormone regulation of glial fibrillar acidic protein immunoreactivity and glial ultrastructure in the rat neuroendocrine hypothalamus. Glia. 1994; 10: 59–60.