

<https://helda.helsinki.fi>

Other Primary Malignancies Among Women With Adult-Type Ovarian Granulosa Cell Tumors

Bryk, Saara

2018-10

Bryk , S , Pukkala , E , Färkkilä , A , Heikinheimo , M , Unkila-Kallio , L & Riska , A 2018 , ' Other Primary Malignancies Among Women With Adult-Type Ovarian Granulosa Cell Tumors ' , International Journal of Gynecological Cancer , vol. 28 , no. 8 , pp. 1529-1534 . <https://doi.org/10.1097/IGC>

<http://hdl.handle.net/10138/311747>

<https://doi.org/10.1097/IGC.0000000000001333>

cc_by_nc

acceptedVersion

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.

This reprint may differ from the original in pagination and typographic detail.

Please cite the original version.

1 **Other Primary Malignancies Among Women With Adult-type Ovarian**
2 **Granulosa Cell Tumors**

3
4
5 Saara Bryk, MD, PhD^{1,2}, Eero Pukkala, PhD^{3,4}, Anniina Färkkilä, MD, PhD^{1,2},
6 Markku Heikinheimo, MD, PhD^{2,5}, Leila Unkila-Kallio, MD, PhD¹, Annika Riska,
7 MD, PhD¹

8
9 1 Obstetrics and Gynecology, University of Helsinki and Helsinki University Hospital

10 2 Children's Hospital, University of Helsinki and Helsinki University Hospital

11 3 Finnish Cancer Registry, Institute for Statistical and Epidemiological Cancer
12 Research, Helsinki, Finland

13 4 Faculty of Social Sciences, University of Tampere, Tampere, Finland

14 5 Department of Pediatrics, Washington University School of Medicine, St Louis
15 Children's Hospital, St Louis, Missouri

16
17 **Corresponding author**

18 Saara Bryk, M.D., Ph.D.

19 Obstetrics and Gynecology, University of Helsinki and Helsinki University Hospital,

20 PO Box 140, 00290 Helsinki, Finland

21 e-mail: saara.bryk@hus.fi

22 Tel: +358503559848

23
24 **Keywords** Granulosa Cell Tumor, Sex Cord-Stromal Tumors, Second Primary

25 Malignancy, Second Primary Cancer, Ovarian Cancer Epidemiology

26
27 **Word count** 2,490

28
29 **Acknowledgments** We thank Tiina Hakanen for invaluable assistance with the cancer
30 registry data. This study received funding from the Finnish Cancer Foundation and
31 University of Helsinki Research Funds.

32
33 The authors declare no conflicts of interest.

34

1 **Abstract**

2

3 **Objective** To determine the incidence of new primary malignancies after adult-type
4 granulosa cell tumor (AGCT), and the incidence of AGCT after breast and uterine
5 cancer using nationwide population-based registry data.

6 **Methods** We identified all patients diagnosed with AGCT in 1968-2013 from the
7 Finnish Cancer Registry (n=986). The number of subsequent primary malignancies
8 among women with AGCT, and the number of AGCTs in women with previous
9 breast or uterine cancer were compared with the expected number of cases, and
10 expressed as Standardized Incidence Ratios (SIRs).

11 **Results** There were 122 cases of subsequent cancers diagnosed at least six months
12 after the primary diagnosis of AGCT (SIR 1.09, 95% CI 0.91-1.3). Particularly, the
13 observed number of cancers of the soft tissue (SIR 4.13, 95% CI 1.33-12.8), thyroid
14 (SIR 3.42, 95% CI 1.54-7.62), and leukemia (SIR 2.67, 95% CI 0.98-5.82) exceeded
15 the number of expected cases. The SIR for breast cancers after AGCT was 1.26 (95%
16 CI 0.92-1.73), and the SIR for AGCT after breast cancer 1.59 (95% CI 1.04-2.29).
17 The risk for subsequent AGCT was more than two-fold in breast cancer patients less
18 than 50 years of age, and over 15 years after primary diagnosis.

19 **Conclusions** There is an increased risk for thyroid and soft tissue cancer as well as
20 leukemia after AGCT, which may be associated with late effects of carcinogenic
21 treatments and possibly shared risk factors. After breast cancer, the risk for AGCT
22 was higher, which may indicate shared hormonal etiology.

23

24

25 **Introduction**

26

27 Adult type-granulosa cell tumors of the ovary (AGCTs) account for 5% of all
28 ovarian neoplasms, and constitute the majority of sex cord-stromal tumors¹. The
29 recently reported age-adjusted (World Standard) incidence rates (truncated to age
30 categories 20 years or older) of AGCT average around 0.6-0.8/100,000 women, and
31 peak after menopause ². AGCTs are characterized by their estrogen-secreting ability,
32 although it has been estimated that approximately 30% of these tumors do not secrete
33 estradiol, likely due to lack of theca cells in the tumor stroma¹. A single somatic point
34 mutation in the transcription factor *FOXL2* (402C-G) is the pathognomonic molecular
35 feature for AGCTs³. Otherwise, the etiological factors remain unknown, although
36 some studies have suggested a possible hormonal background for these tumors^{4, 5}.
37 According to current knowledge, there is no hereditary predisposition for the
38 development of AGCT.

39 The diagnosis is usually made at an early stage, partly due to symptoms
40 related to hormone secretion, and the disease tends to run an indolent course.
41 Excessive exposure to tumor-derived estrogen among these patients leads to an
42 increased risk of concomitant endometrial pathology and endometrial cancer ⁶⁻⁹.
43 There are, however, only a few studies focusing on other primary malignancies in
44 women with AGCT ^{10, 11}. In general, the risk of other primary malignancies after
45 ovarian cancer is associated with either inherent genetic or lifestyle-related extrinsic
46 risk factors, or carcinogenic treatment regimens ¹²⁻¹⁴. As other, particularly endocrine-
47 related cancers may share etiological factors with AGCTs, it is of interest to study the
48 potential association of these cancers, especially breast and uterine cancer. The object

49 of our study was to evaluate the incidence of all other primary cancers after AGCT, as
50 well the incidence of AGCT after breast or uterine cancer.

51

52 **Materials and methods**

53

54 In this retrospective cohort study, we identified all patients diagnosed with
55 AGCT in Finland during 1968-2013 from the Finnish Cancer Registry (FCR). The
56 FCR is a high-quality, population-based registry relying on unique personal identity
57 codes. The personal identity code is a specific means of identification, which remains
58 unchanged throughout the person's lifetime, and has been used in Finland since the
59 1960s. Physicians, hospitals, and pathology and hematology laboratories in Finland
60 are obliged to report all malignant tumors to the FCR, resulting in a nearly complete
61 registration of all cancer cases¹⁵. Information on vital status and emigration was
62 obtained from the Population Register Center, which is directly linked to the FCR
63 information.

64 AGCTs were retrieved from the registry applying the ICD topography code
65 C56.9 with morphology codes M8620/1, 8620/3, 8621/1, and 8621/3. The incidence
66 rates of AGCT during the follow-up period were calculated, and adjusted for age to
67 the World Standard Population. All patients were followed up for second primary
68 cancer from the date of first diagnosis (1968-2013) to the date of death, date of
69 migration, or until December 31st, 2013. In order to identify concomitant cancers and
70 surveillance bias, the analyses were carried out in two subgroups: 1) all subsequent
71 tumors after AGCT, and 2) all subsequent tumors except those occurring within six
72 months after AGCT. Subsequent primary tumors were grouped in 18 categories based
73 on cancer site, and included ICD-codes C00-96, D32-33, D42-43, D45-47, and D76

74 (mouth/pharynx, digestive organs, respiratory organs, breast, female genitalia, urinary
75 organs, melanoma of the skin, skin (other than melanoma), eye, thyroid gland, other
76 endocrine glands, bone, soft tissues, mesothelioma, autonomic nervous system,
77 brain/central nervous system, lymphoid/hematopoietic tissue, other/not defined). The
78 number of new primary malignant tumors among women with previous AGCT was
79 compared with the expected number of cases calculated from the accumulated person-
80 years and incidence rates for the national population, stratified by age and year of
81 diagnosis.

82 Secondly, we analyzed the number of subsequent AGCTs in women with a
83 first primary breast or uterine cancer (ICD C54 and C50), and compared it with the
84 expected number of AGCTs. These analyses were likewise performed separately on
85 all subsequent AGCTs as well as those occurring within 6 months of the primary
86 cancer diagnoses. The ratio of observed to expected cases was defined as the
87 Standardized Incidence Ratio (SIR), and 95% confidence intervals (CI) were
88 calculated. The SIRs were also stratified for time since first primary cancer diagnosis
89 (0-4 years, 5-14 and 15+ years after the diagnosis of first primary tumor), for age at
90 the first primary cancer diagnosis (<50 or 50 years or older), and in breast cancer also
91 for the invasion status (localized vs. non-localized).

92 The ethics committee of Helsinki University Hospital (HUH) and the National
93 Supervisory Authority for Welfare and Health approved the study.

94

95

96 **Results**

97

98 In 1968-2013, a total of 986 women in Finland were diagnosed with AGCT.
99 The age-adjusted (World Standard) incidence varied between 0.4 and 0.9 per 100,000
100 women, with approximately 20 cases each year (Figure 1). The logarithmic trend line
101 suggests a decreasing trend in the incidence of AGCT over the 45-year study period.

102 After the diagnosis of AGCT, 122 cases of new primary malignant tumors
103 were recorded, resulting in a 12.4% rate of second malignancies among AGCT
104 patients. The expected number was 111.7 (SIR 1.09, 95% CI 0.91-1.3) (Table 1). If
105 also cancers diagnosed within six months of AGCT were included, the total rate was
106 13.9% and SIR 1.19 (95% CI 1-1.41, $p=0.04$). The SIR for these cancers only was
107 5.00 (95% CI 2.80-8.23). The median interval between the diagnosis of AGCT and
108 second primary tumor was 19.2 years (range 0.02-45.6 years). In a minimum time of
109 six months from the primary cancer diagnosis, the observed number of thyroid cancer,
110 soft tissue cancer, and leukemia exceeded the number of expected cases significantly
111 (Table 1). The SIRs were also elevated for cancers of the oropharynx, breast, urinary
112 organs, skin (non-melanoma), and mesothelioma, but not significantly (Table 1).
113 There were less than expected cancer cases in the uterine corpus and ovaries. For all
114 subsequent cancers, the SIRs stratified for follow-up time were 0.75 for 0.5-4 years
115 (95% CI 0.45-1.15), 0.98 for 5-14 years (95% CI 0.72-1.30), and 1.40 (95% CI 1.07-
116 1.78) for more than 15 years after diagnosis of AGCT. The SIR was higher for
117 patients who were less than 50 years of age at primary diagnosis (SIR 1.31, 95% CI
118 0.96-1.75). The results were largely similar when also second primary cancer cases
119 diagnosed within six months after AGCT were included, with the exception of uterine
120 cancer.

121 The SIR for breast cancer after AGCT was 1.4 after at least five years of
122 primary cancer diagnosis (Table 2). Subsequent breast cancer was somewhat more

123 common in patients who were at least 50 years old at the time of AGCT diagnosis
124 (SIR 1.31, 95% CI 0.86-1.91). The SIR was only elevated in localized breast cancer
125 (SIR 1.36, 95% CI 0.86-2.02), as opposed to non-localized breast cancer (SIR 0.83,
126 95% CI 0.42-1.46). In patients who had breast cancer diagnosed primarily, there were
127 25 cases of subsequent AGCTs during follow-up (Table 3). The SIR for AGCT after
128 breast cancer was 1.59 (95% CI 1.04-2.29), and increased with time since breast
129 cancer diagnosis to 2.28 (95% CI 0.98-4.41) in the follow-up category of 15 years or
130 more. For age below 50 years at breast cancer diagnosis the SIR was 2.10 (95 % CI
131 1.09-3.59).

132 From the cancers diagnosed within six months of AGCT, uterine cancer
133 accounted for 33% (n=5), digestive organs 27% (n=4), and breast cancer 13% (n=2)
134 of these cases. Other malignancies reported within this follow-up period included
135 cancers of the urinary tract, and lymphoid/hematopoietic tissue. After uterine cancer,
136 AGCT was diagnosed in 20 women within six months (SIR 4.99, 95% CI 3.18-7.37),
137 whereas two women developed AGCT more than 6 months after primary uterine
138 cancer diagnosis. All women with uterine cancer and subsequent AGCT were at least
139 50 years of age at the time of the uterine cancer diagnosis (SIR 6.21, 95% CI 4.09-
140 9.42).

141

142 **Discussion**

143

144 The indolent course, relatively low disease-related mortality and estrogen-
145 secreting capability of AGCT result in a clinically relevant lifetime risk for
146 developing a second primary cancer. On the other hand, the etiological factors of
147 AGCT are largely unknown, and a common predisposing factor may exist behind

148 AGCT and other hormone-related cancers. To our knowledge, this is the largest and
149 first study since Björkholm et al. in 1980¹¹ to analyze all second primary malignancies
150 among AGCT patients. In our study, women with AGCT had a 9% increased risk of
151 developing a new primary malignancy as compared with the general population. If
152 cancers diagnosed within six months after the primary tumor were included, the risk
153 was significantly increased by 19%. The large difference in these figures is mainly
154 explained by the presence of concomitant endometrial cancer, but the significant
155 number of cancers of the digestive organs diagnosed within six months of AGCT
156 most likely also reflects the increased surveillance among cancer patients in general.

157 Two recent publications have described the incidence of endometrial cancer
158 and breast cancer among patients with AGCT^{6, 10}. Van Meurs et al. found a 6% rate
159 of endometrial cancer concomitant with the diagnosis of AGCT, but no increased risk
160 for endometrial abnormalities in the median follow-up time of 10 years after AGCT
161 for patients not having undergone hysterectomy⁶. Other population-based studies have
162 reported 5-8% rates of concomitant endometrial cancer^{8, 9, 11}, and we reported similar
163 rates in a large, single-institute patient cohort¹⁶. In the current population-based
164 registry cohort, the rate was 2.5% when patients diagnosed primarily with either
165 AGCT or uterine cancer and a subsequent uterine cancer or AGCT within six months
166 of primary diagnosis were included. This relatively low rate may reflect a proportion
167 of previously hysterectomized patients, since they could not be excluded from the
168 original cancer registry data. This would also explain the higher incidence in hospital-
169 based cohorts, as solely patients with endometrial sample available have been
170 evaluated.

171 We found an increased risk for breast cancer both before and after diagnosis of
172 AGCT, although the risk was significant only before AGCT. This is a similar finding

173 to the smaller Danish, Israeli and US studies where the rate of breast cancer among
174 AGCT patients was 5-10% ^{9, 10, 17}. In our study, the rate of breast cancer was 6.9%
175 among all women with AGCT. After AGCT, the risk was confined to localized breast
176 cancers, which may indicate towards surveillance bias, i.e. the increased frequency
177 and intensity of clinical follow-up and examination among patients with previously
178 diagnosed cancer. There was a relatively long latency between breast cancer and
179 AGCT regardless of which cancer was the first primary tumor, which does not
180 support genetic susceptibility. AGCT is neither associated with any of the known
181 predisposing mutations to breast cancer such as BRCA1 and BRCA2 mutations, nor is
182 the FOXL2 mutation pathognomonic to AGCT present in breast carcinoma ^{1, 18}. In
183 the present study, the risk for subsequent AGCT in breast cancer patients was
184 significantly increased in women who were younger than 50 years at primary
185 diagnosis, which probably reflects the long follow-up time, as the latency between the
186 cancers was also long. Shared etiological factors such as obesity, parity, and hormonal
187 environment offer a possible explanation for the increased incidence of breast cancer
188 and AGCT among same women. Obesity represents a hyperestrogenic state and is a
189 known risk factor for breast cancer¹⁹, and has been suggested as a risk factor for
190 AGCT⁴. In post-menopausal women, breast cancer risk is around twice as high in
191 those with the highest sex hormone levels compared to those with the lowest²⁰. Parity,
192 on the other hand, is a protective factor in both breast and ovarian cancer ^{21, 22}.

193 The effects of primary cancer treatment may influence the development of
194 second primary AGCT. Selective estrogen receptor modulators (SERMs) such as
195 tamoxifen are used to treat hormone-receptor positive breast cancer, and three case
196 reports have linked antecedent tamoxifen use with the development of AGCT²³⁻²⁵.
197 Furthermore, aromatase inhibitors such as letrozole are used in the treatment of both

198 postmenopausal breast cancer and AGCT^{26, 27}. However, further evidence is
199 warranted to establish a causal link between hormonal breast cancer treatment and the
200 development of AGCT.

201 Similarly to an earlier study, we found an increased risk for thyroid cancer
202 among patients with previous AGCT, but it should be noted that the number of cases
203 is rather small in both studies¹¹. It has been proposed that female hormones,
204 reproductive factors, and obesity also play a role in thyroid cancer pathogenesis, but
205 there are no consistent data linking ovarian and thyroid cancer^{28, 29}. *DICER1* germline
206 mutation carriers have a predisposition to both thyroid cancer and sex cord-stromal,
207 particularly Sertoli-Leydig cell tumors^{30, 31}. Cancer registry data are not, however,
208 molecularly validated, and there is a possibility that some of the tumors identified as
209 AGCT may actually represent other sex cord-stromal tumors.

210 We also found significantly increased SIRs for soft tissue cancer and leukemia
211 after AGCT. The development of secondary soft tissue sarcoma is strongly associated
212 with radiation exposure from radiotherapy, especially after breast cancer^{32, 33}. No
213 association between female reproductive factors and the development of soft tissue
214 cancer has been detected, but the studies in this field are scarce³⁴. As nowadays
215 radiotherapy is rarely used in the treatment of AGCT, the increased SIR for soft tissue
216 cancer is most likely related to shared risk factors. Furthermore, radiotherapy for
217 ovarian cancer is known to be associated with bladder carcinoma, but in our series,
218 the SIR for bladder cancer was not significantly elevated after AGCT³⁵. Adjuvant
219 therapy is used in the management of metastatic or recurrent AGCT, and presently
220 consists of platinum-based chemotherapeutic agents or more recently, hormonal
221 treatments^{26, 36}. Late effects of chemotherapy may include increased risk for leukemia,
222 and most likely explains the high incidence of this cancer among patients with

223 previous AGCT^{12, 14}. Two large population-based studies on second malignancies
224 after ovarian cancer of any type both reported significantly elevated SIRs for cancers
225 of the colon, rectum, lung, breast, bladder, and thyroid, as well as for leukemia, and
226 the risk for subsequent cancer development was associated with older age, chemo-
227 and radiotherapy^{12, 35}.

228 This is the largest study to date in analyzing the risk for other primary
229 malignancies associated with AGCT. The strengths of this study are the reliable and
230 comprehensive cancer registry incidence data, and the long observation period of over
231 40 years. The rarity and the lack of molecular validation of AGCT, as well as of
232 individual data such as parity, BMI, or use of hormonal therapies are limiting factors
233 in this analysis.

234 In conclusion, we found a slightly elevated risk for overall second
235 malignancy, particularly thyroid and soft tissue cancer, and leukemia. Partly these
236 excesses may result from carcinogenic treatments for AGCT. The increased incidence
237 of AGCT and breast cancer among the very same patients may indicate shared
238 hormonal etiology. Earlier studies have concluded that breast cancer patients have a
239 higher incidence of second primary ovarian cancer, particularly when diagnosed
240 before 50 years of age; this patient group might benefit from regular gynecological
241 surveillance³⁷⁻³⁹. This seems to be true also for AGCT after breast cancer, which
242 should be recognized in patient counseling and long-term clinical follow-up after the
243 primary tumor.

244

245

246

247

248

249

250 **Acknowledgements** We thank Tiina Hakanen for invaluable assistance with the
251 cancer registry data. This study received funding from the Finnish Cancer Foundation
252 and University of Helsinki Research Funds.

253

254 The authors declare no conflict of interest.

255

256

257

258 **References**

259

- 260 1. Schumer ST, Cannistra SA. Granulosa cell tumor of the ovary. *Journal of*
261 *clinical oncology : official journal of the American Society of Clinical Oncology.*
262 2003;21(6):1180-9.
- 263 2. Bryk S, Pukkala E, Martinsen JI, et al. Incidence and occupational variation
264 of ovarian granulosa cell tumours in Finland, Iceland, Norway and Sweden
265 during 1953-2012: a longitudinal cohort study. *BJOG.* 2017;124(1):143-9.
- 266 3. Jamieson S, Butzow R, Andersson N, et al. The FOXL2 C134W mutation is
267 characteristic of adult granulosa cell tumors of the ovary. *Modern pathology : an*
268 *official journal of the United States and Canadian Academy of Pathology, Inc.*
269 2010;23(11):1477-85.
- 270 4. Boyce EA, Costaggini I, Vitonis A, et al. The epidemiology of ovarian
271 granulosa cell tumors: a case-control study. *Gynecol Oncol.* 2009;115(2):221-5.
- 272 5. Chen T, Surcel HM, Lundin E, et al. Circulating sex steroids during
273 pregnancy and maternal risk of non-epithelial ovarian cancer. *Cancer*
274 *epidemiology, biomarkers & prevention : a publication of the American*
275 *Association for Cancer Research, cosponsored by the American Society of*
276 *Preventive Oncology.* 2011;20(2):324-36.
- 277 6. van Meurs HS, Bleeker MC, van der Velden J, et al. The incidence of
278 endometrial hyperplasia and cancer in 1031 patients with a granulosa cell tumor
279 of the ovary: long-term follow-up in a population-based cohort study. *Int J*
280 *Gynecol Cancer.* 2013;23(8):1417-22.
- 281 7. Ayhan A, Salman MC, Velipasaoglu M, et al. Prognostic factors in adult
282 granulosa cell tumors of the ovary: a retrospective analysis of 80 cases. *J Gynecol*
283 *Oncol.* 2009;20(3):158-63.
- 284 8. Unkila-Kallio L, Tiitinen A, Wahlström T, et al. Reproductive features in
285 women developing ovarian granulosa cell tumour at a fertile age. *Hum Reprod.*
286 2000;15(3):589-93.
- 287 9. Ohel G, Kaneti H, Schenker JG. Granulosa cell tumors in Israel: a study of
288 172 cases. *Gynecol Oncol.* 1983;15(2):278-86.
- 289 10. Hammer A, Lauszus FF, Petersen AC. Ovarian granulosa cell tumor and
290 increased risk of breast cancer. *Acta Obstet Gynecol Scand.* 2013;92(12):1422-5.
- 291 11. Björkholm E, Silfverswärd C. Granulosa- and theca-cell tumors. Incidence
292 and occurrence of second primary tumors. *Acta Radiol Oncol.* 1980;19(3):161-7.
- 293 12. Hung YP, Liu CJ, Hu YW, et al. Secondary Primary Malignancy Risk in
294 Patients With Ovarian Cancer in Taiwan: A Nationwide Population-Based Study.
295 *Medicine.* 2015;94(38):e1626.
- 296 13. Keum N, Greenwood DC, Lee DH, et al. Adult weight gain and adiposity-
297 related cancers: a dose-response meta-analysis of prospective observational
298 studies. *Journal of the National Cancer Institute.* 2015;107(2).
- 299 14. Travis LB, Holowaty EJ, Bergfeldt K, et al. Risk of leukemia after platinum-
300 based chemotherapy for ovarian cancer. *N Engl J Med.* 1999;340(5):351-7.
- 301 15. Pukkala E, Engholm G, Hojsgaard Schmidt LK, et al. Nordic Cancer
302 Registries - an overview of their procedures and data comparability. *Acta Oncol.*
303 2017:1-16.

- 304 16. Bryk S, Farkkila A, Butzow R, et al. Clinical characteristics and survival of
305 patients with an adult-type ovarian granulosa cell tumor: a 56-year single-center
306 experience. *Int J Gynecol Cancer*. 2015;25(1):33-41.
- 307 17. Meisel JL, Hyman DM, Jotwani A, et al. The role of systemic chemotherapy
308 in the management of granulosa cell tumors. *Gynecol Oncol*. 2015;136(3):505-
309 11.
- 310 18. Shah SP, K̄abel M, Senz J, et al. Mutation of FOXL2 in granulosa-cell
311 tumors of the ovary. *N Engl J Med*. 2009;360(26):2719-29.
- 312 19. Kyrgiou M, Kalliala I, Markozannes G, et al. Adiposity and cancer at major
313 anatomical sites: umbrella review of the literature. *BMJ*. 2017;356:j477.
- 314 20. Key T, Appleby P, Barnes I, et al. Endogenous sex hormones and breast
315 cancer in postmenopausal women: reanalysis of nine prospective studies.
316 *Journal of the National Cancer Institute*. 2002;94(8):606-16.
- 317 21. Adami HO, Hsieh CC, Lambe M, et al. Parity, age at first childbirth, and risk
318 of ovarian cancer. *Lancet*. 1994;344(8932):1250-4.
- 319 22. Layde PM, Webster LA, Baughman AL, et al. The independent associations
320 of parity, age at first full term pregnancy, and duration of breastfeeding with the
321 risk of breast cancer. Cancer and Steroid Hormone Study Group. *Journal of
322 clinical epidemiology*. 1989;42(10):963-73.
- 323 23. Gherman RB, Parker MF, Macri CI. Granulosa cell tumor of the ovary
324 associated with antecedent tamoxifen use. *Obstetrics and gynecology*. 1994;84(4
325 Pt 2):717-9.
- 326 24. Abahssain H, Kairouani M, Gherman R, et al. Granulosa cell tumor of the
327 ovary and antecedent of adjuvant tamoxifen use for breast cancer. *World journal
328 of surgical oncology*. 2010;8:67.
- 329 25. Tanaka T, Kato T, Ohmichi M. Granulosa cell tumor of the ovary after long-
330 term use of tamoxifen and toremifene. *The journal of obstetrics and gynaecology
331 research*. 2012;38(12):1379-84.
- 332 26. van Meurs HS, van Lonkhuijzen LR, Limpens J, et al. Hormone therapy in
333 ovarian granulosa cell tumors: a systematic review. *Gynecol Oncol*.
334 2014;134(1):196-205.
- 335 27. Schiavon G, Smith IE. Status of adjuvant endocrine therapy for breast
336 cancer. *Breast cancer research : BCR*. 2014;16(2):206.
- 337 28. Negri E, Dal Maso L, Ron E, et al. A pooled analysis of case-control studies
338 of thyroid cancer. II. Menstrual and reproductive factors. *Cancer Causes Control*.
339 1999;10(2):143-55.
- 340 29. Zhu J, Zhu X, Tu C, et al. Parity and thyroid cancer risk: a meta-analysis of
341 epidemiological studies. *Cancer Med*. 2016;5(4):739-52.
- 342 30. Rutter MM, Jha P, Schultz KA, et al. DICER1 Mutations and Differentiated
343 Thyroid Carcinoma: Evidence of a Direct Association. *J Clin Endocrinol Metab*.
344 2016;101(1):1-5.
- 345 31. Stewart CJ, Charles A, Foulkes WD. Gynecologic Manifestations of the
346 DICER1 Syndrome. *Surgical pathology clinics*. 2016;9(2):227-41.
- 347 32. Fletcher CD. The evolving classification of soft tissue tumours - an update
348 based on the new 2013 WHO classification. *Histopathology*. 2014;64(1):2-11.
- 349 33. Virtanen A, Pukkala E, Auvinen A. Incidence of bone and soft tissue
350 sarcoma after radiotherapy: a cohort study of 295,712 Finnish cancer patients.
351 *Int J Cancer*. 2006;118(4):1017-21.

352 34. Fioretti F, Tavani A, Gallus S, et al. Menstrual and reproductive factors and
353 risk of soft tissue sarcomas. *Cancer*. 2000;88(4):786-9.

354 35. Travis LB, Curtis RE, Boice JD, Jr., et al. Second malignant neoplasms
355 among long-term survivors of ovarian cancer. *Cancer research*.
356 1996;56(7):1564-70.

357 36. Savage P, Constenla D, Fisher C, et al. Granulosa cell tumours of the ovary:
358 demographics, survival and the management of advanced disease. *Clinical*
359 *oncology*. 1998;10(4):242-5.

360 37. Molina-Montes E, Pollan M, Payer T, et al. Risk of second primary cancer
361 among women with breast cancer: a population-based study in Granada (Spain).
362 *Gynecol Oncol*. 2013;130(2):340-5.

363 38. Kirova YM, De Rycke Y, Gambotti L, et al. Second malignancies after breast
364 cancer: the impact of different treatment modalities. *Br J Cancer*.
365 2008;98(5):870-4.

366 39. Hall HI, Jamison P, Weir HK. Second primary ovarian cancer among
367 women diagnosed previously with cancer. *Cancer epidemiology, biomarkers &*
368 *prevention : a publication of the American Association for Cancer Research,*
369 *cosponsored by the American Society of Preventive Oncology*. 2001;10(9):995-9.
370

371

372

373

374

375

376 **Figure legends**

377

378 Figure 1. Incidence of adult-type ovarian granulosa cell tumors (AGCTs) in Finland
379 in 1968-2013, with a logarithmic trend line (dotted).

380

381

382

383

384

Table 1. Risk of subsequent primary malignancies among Finnish women with previous adult-type ovarian granulosa cell tumor (AGCT) in 1968-2013, by site.

Second primary tumor site	Observed	Expected	SIR	95% CI	p-value
All	137	114.7	1.19	1-1.41	0.04
All, diagnosis within 6 months of AGCT	15	3.00	5.00	2.80-8.23	<0.001
All, diagnosis > 6 months after AGCT	122	111.7	1.09	0.91-1.3	0.33
Mouth, pharynx	3	1.7	1.76	0.57-5.45	0.33
Digestive organs	24	25.7	0.93	0.63-1.39	0.74
Respiratory organs	5	6.3	0.79	0.33-1.90	0.60
Skin, melanoma	3	3.0	0.99	0.32-3.06	0.98
Skin, non-melanoma	9	4.8	1.86	0.97-3.58	0.06
Soft tissues	3	0.7	4.13	1.33-12.8	0.01
Breast	38	30.2	1.26	0.92-1.73	0.15
Female genitalia	5	14.3	0.35	0.15-0.84	0.02
Corpus uteri	0	7.1	0.00	0.00-0.52	0.01
Ovary	1	4.3	0.23	0.01-1.30	0.18
Cervix uteri	2	1.3	1.49	0.18-5.38	0.89
Other	2	1.5	1.38	0.17-4.98	0.97
Urinary organs	8	5.6	1.43	0.71-2.86	0.31
Bladder and urinary tract	3	2.4	1.27	0.26-3.73	0.92
Brain, central nervous system	3	3.9	0.76	0.25-2.36	0.64
Thyroid gland	6	1.8	3.42	1.54-7.62	0.003
Lymphoid and hematopoietic tissue	11	9.6	1.15	0.64-2.07	0.65
Leukemia	6	2.2	2.67	0.98-5.82	0.03
Other or not defined	3	3.4	0.88	0.28-2.72	0.82

SIR= standardized incidence ratio, CI = confidence interval. Sites with < 3 observed cases are excluded, with the exception of cancers of the female genitalia.

Table 2. Risk of subsequent breast cancer among Finnish women with previous adult-type granulosa cell tumor (AGCT) in 1968-2013, by age at and time since AGCT diagnosis, and breast cancer invasion.

	Observed	Expected	SIR	95% CI	p-value
All	40	31	1.29	0.93-1.73	0.11
All, diagnosis > 6 months after AGCT	38	30.2	1.26	0.9-1.7	0.15
Follow-up time (years)					
0-4	6	6.9	0.87	0.34-1.76	0.73
5-14	17	12.3	1.38	0.82-2.15	0.18
≥15	15	10.9	1.37	0.79-2.19	0.22
Age at AGCT diagnosis					
<50	14	11.9	1.18	0.66-1.91	0.54
≥50	24	18.3	1.31	0.86-1.91	0.18
Breast cancer invasion ¹					
Localized	21	15.5	1.36	0.86-2.02	0.16
Non-localized	10	12.0	0.83	0.42-1.46	0.56

AGCT= adult-type ovarian granulosa cell tumor, SIR= standardized incidence ratio, CI = confidence interval.

¹Invasion status unknown in seven cases

Table 3. Risk of subsequent adult-type granulosa cell tumors (AGCTs) among Finnish women with breast cancer in 1968-2013, by age at and time since breast cancer diagnosis.

	Observed	Expected	SIR	95% CI	p-value
All	28	16.6	1.69	1.14-2.4	0.006
All, diagnosis > 6 months after breast cancer	25	15.7	1.59	1.04-2.29	0.02
Follow-up time (years)					
0-4	8	5.9	1.35	0.62-2.52	0.39
5-14	10	6.8	1.48	0.74-2.59	0.22
≥15	7	3.1	2.28	0.98-4.41	0.03
Age at breast cancer diagnosis					
<50	11	5.2	2.10	1.09-3.59	0.01
≥50	14	10.5	1.33	0.75-2.16	0.28

SIR= standardized incidence ratio, CI = confidence interval.

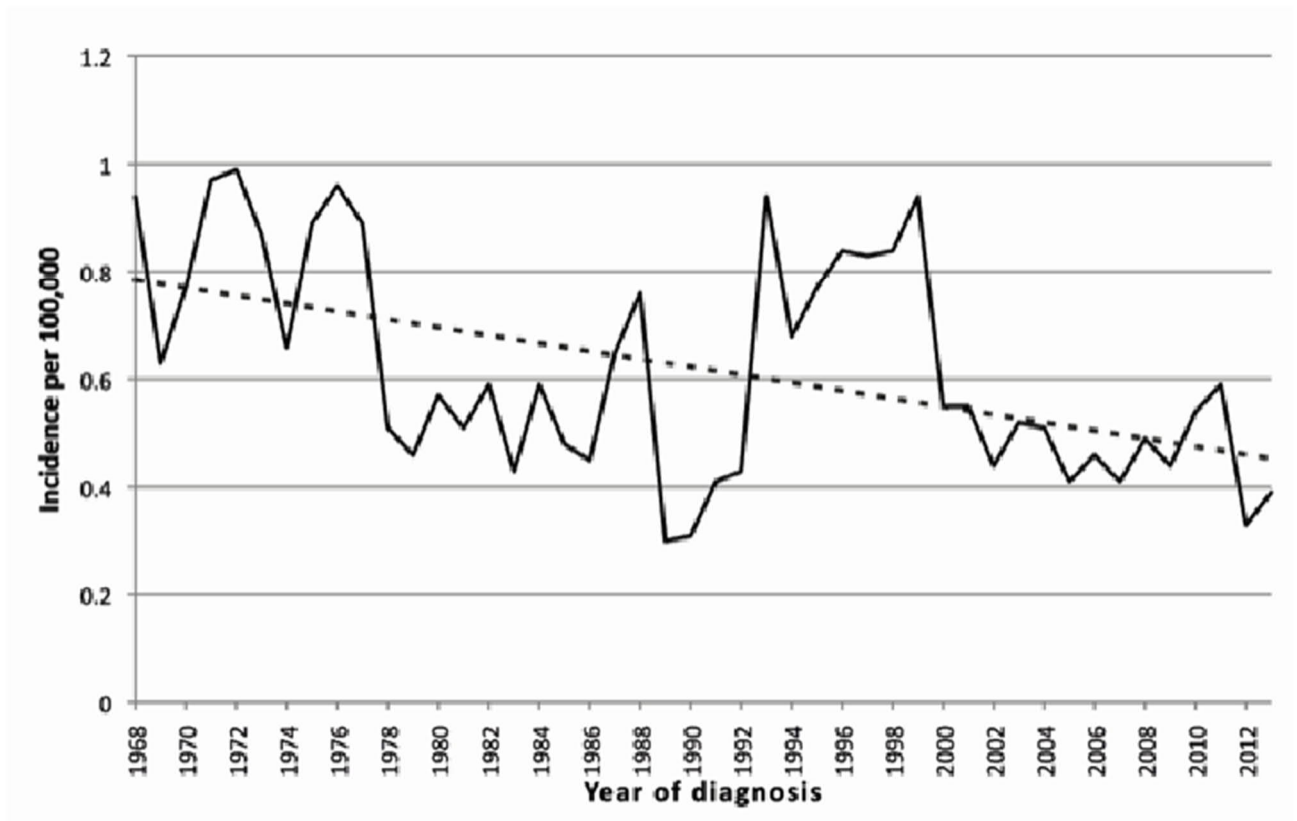


Figure 1. Incidence of adult-type ovarian granulosa cell tumors (AGCTs) in Finland in 1968-2013, with a logarithmic trend line (dotted).