Background: Infertility may indicate an underlying predisposition toward premature cardiovascular disease, yet little is known about potential long-term cardiovascular events following fertility therapy. We investigated whether failure of fertility therapy is associated with subsequent adverse cardiovascular events.

Methods: We performed a population-based cohort analysis of women who received gonadotropin-based fertility therapy between Apr. 1, 1993, and Mar. 31, 2011, distinguishing those who subsequently gave birth and those who did not. Using multivariable Poisson regression models, we estimated the relative rate ratio of adverse cardiovascular events associated with fertility therapy failure, accounting for age, year, baseline risk factors, health care history and number of fertility cycles. The primary outcome was subsequent treatment for nonfatal coronary ischemia, stroke, transient ischemic attack, heart failure or thromboembolism.

Results: Of 28,442 women who received fertility therapy, 9,349 (32.9%) subsequently gave birth and 19,093 (67.1%) did not. The median number of fertility treatments was 3 (interquartile range 1–5). We identified 2,686 cardiovascular events over a median 8.4 years of follow-up. The annual rate of cardiovascular events was 19% higher among women who did not give birth after fertility therapy than among those who did (1.08 v. 0.91 per 100 patient-years, p < 0.001), equivalent to a 21% relative increase in the annual rate (95% confidence interval 13%–30%). We observed no association between event rates and number of treatment cycles.

Interpretation: Fertility therapy failure was associated with an increased risk of long-term adverse cardiovascular events. These women merit surveillance for subsequent cardiovascular events.

Failure of fertility therapy and subsequent adverse cardiovascular events

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ABSTRACT

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methods to screen for a subsequent newborn delivery (OHIP codes P006, P018 and P020). Overall, the OHIP databases have a high level of completeness (> 99%) and diagnostic accuracy (> 95%) in this setting.24,25 Fertility therapy failure was defined as cycles of treatment that were not followed by a subsequent newborn delivery within 1 year. Because individuals may have undergone repeated cycles of fertility therapy, we classified each woman by the total number of cycles of treatment by analyzing consecutive 28-day blocks to explore potential dose–response relationships.

The available databases lacked information on specific fertility medications because Ontario did not provide single-payer universal coverage for fertility prescriptions. However, human menopausal gonadotropins and gonadotropin-releasing hormone both qualified for OHIP code G334.

**Patient characteristics**

We collected baseline data on demographic and clinical characteristics from the Canadian Institute for Health Information hospital and outpatient databases during the 2 years before each woman’s final fertility treatment.28 The databases also served as the source for identifying follow-up outcomes, admissions and procedures. In addition, we used the hospital database to collect information on the patient’s age and sex, date of admission, and additional diagnoses coded with the International Classification of Diseases, 9th revision (ICD-9; 16 diagnoses) or the International Statistical Classification of Diseases and Related Health Problems, 10th revision (ICD-10; 25 diagnoses).

Some diagnoses were primarily identified outside of hospital (e.g., hypertension and diabetes); we used the outpatient database to determine selected diagnoses and aggregate health care utilization accumulated during the 2 years before the final fertility treatment.30,31 We estimated the patient’s residence location (urban or rural) and neighbourhood income (in quintiles) from the individual’s home address. History of live births, miscarriages and stillbirths was ascertained over the duration of available data. We also collected information on history of hypertension, diabetes, hypercholesterolemia, smoking, obesity, gestational hypertension, gestational diabetes, ischemic heart disease, heart failure, atrial fibrillation, asthma, neoplasm and depression, and number of health care visits in the prior 2 years, number of fertility cycles and number of in-vitro fertilization procedures. Information about laboratory test results and prescription medications were not available.

**Outcomes**

The primary outcome was a composite of medical care for nonfatal coronary ischemia, acute stroke, transient ischemic attack (TIA), heart failure or thromboembolism. We used ICD-9 codes to identify study outcomes before Mar. 31, 2002, and ICD-10 codes to identify those diagnosed from Apr. 1, 2002, onward to account for changes in databases over time (Appendix 1, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.160744/-/DC1). Prior validation studies suggested that these codes have mid-range sensitivity (58% to 89% for coronary ischemia and heart failure), excellent specificity (93% to 97% for coronary ischemia and heart failure) and a reasonable positive predictive value (65% to 95% for thromboembolism, and 90% to 92% for stroke or TIA).32–34 Secondary outcomes were the individual components of the primary outcome.

Cardiovascular mortality was not included in the primary outcome measure because of the rarity of the event (total mortality was examined in sensitivity analyses).

**Statistical analysis**

We compared baseline demographic and clinical characteristics between groups using the χ² test for categorical variables and the t test for continuous variables. We estimated annualized event rates for the primary and secondary outcomes by dividing the total number of events by the total number of person-years of follow-up. We followed all patients until study termination (Mar. 31, 2015) and included both first and repeat events. We used Poisson regression models with generalized estimating equations to estimate the relative rate ratio and 95% confidence interval (CI) of cardiovascular events associated with fertility therapy failure, with time zero set at 1 year following the final fertility treatment.

In multivariable analyses, we adjusted for potential confounders, including age (in years), calendar year, geographic residence (rural v. urban), neighbourhood income (in quintiles), prior health care visits, cardiovascular risk factors (hypertension, diabetes, hypercholesterolemia, smoking and obesity), ischemic heart disease, heart failure, atrial fibrillation, asthma, depression, neoplasm, prior live births, miscarriages and stillbirths, and number of fertility cycles. We performed tests for departure from linearity for continuous variables before estimating a linear association. For patients without recorded diagnoses, we included a code indicating no disease so that all patients could be included in all analyses. We further explored potential effect modification by introducing interaction terms assessing fertility therapy failure with age, type of infertility (primary or secondary) and history of miscarriage. We tested whether a dose–response relationship was present by analyzing results according to whether patients received a single cycle or multiple cycles of fertility therapy.

We conducted survival analyses restricted to the time to first hospital admission for a primary outcome or death using a Cox proportional hazards model, with age as the underlying time variable to derive hazards ratios (HRs) and to check the robustness of the primary analysis. Potential departures from the proportionality assumption were evaluated with the introduction of a time-varying effect term for fertility therapy failure into the multivariable adjusted Cox model.

All p values were 2-sided, and a threshold less than 0.05 was considered significant. All statistical analyses were performed with the use of SAS version 9.3 (SAS Institute).

**Ethics approval**

The Research Ethics Board of Sunnybrook Health Sciences Centre approved the study protocol.

**Results**

A total of 28 442 women received fertility therapy during the study, of whom 9349 (32.9%) gave birth and 19 093 (67.1%) did not give birth within 1 year after the final treatment (Table 1). The mean age was
35.2 (standard deviation 4.8) years, and 23575 (82.9%) had no prior obstetric deliveries. The median number of fertility cycles was 3 (interquartile range 1–5); 7649 (26.9%) of the patients received a single cycle of treatment, and 20793 (73.1%) received multiple cycles.

Fertility therapy failure was most common among women who were older, were living in lower income neighbourhoods, used less health care and had fewer miscarriages or stillbirths. Diabetes and hypertension (including gestational presentations) were the 2 most common traditional cardiovascular risk factors, and each was less common among women who did not give birth following fertility therapy than among those who did (p < 0.001). In contrast, hypercholesterolemia (p < 0.001), smoking (p = 0.02), asthma (p = 0.02), neoplasm (p < 0.001) and depression (p = 0.03) were more common among women who did not give birth following fertility therapy.

During a median follow-up of 8.4 years (interquartile range 5.3–12.4), a total of 2686 primary cardiovascular events were observed among women who underwent fertility therapy, of which 1925 occurred among women who did not subsequently give birth and 761 occurred among those who gave birth. The annual rate of cardiovascular events was 19% higher among women who did not give birth than among those who did following fertility therapy (1.08 v. 0.91 events per 100 patient-years; relative rate ratio 1.19, 95% CI 1.11–1.27). The observed increase persisted after multivariable adjustment (adjusted relative rate ratio 1.21, 95% CI 1.13–1.30). Among the specific components of the primary outcome, a total of 695 outpatient visits were for cerebrovascular events (624 ischemic strokes or TIAs, 71 hemorrhagic strokes), 940 for ven thromboembolism, 567 for coronary ischemia and 484 for heart failure. The overall increase in the cardiovascular event rate was explained mostly by increases in the rate of heart failure (adjusted relative rate ratio 2.25, 95% CI 2.06–2.46) and cerebrovascular events (adjusted relative rate ratio 1.25, 95% CI 1.15–1.37), with an adjusted relative rate ratio of 1.08 for ven thromboembolic events (95% CI 1.00–1.17). In contrast, no increase was observed between fertility therapy failure and coronary ischemia (Table 2). When cerebrovascular risk was further subcategorized, fertility therapy failure was associated with an increased risk of ischemic stroke (adjusted relative rate ratio 1.33, 95% CI 1.22–1.46) but not hemorrhagic stroke (adjusted relative rate ratio 0.88, 95% CI 0.80–0.96).

In the survival analysis restricted to the time to first hospital admission for the composite outcome of a cardiovascular endpoint or death, the risk was increased among women who had fertility therapy failure compared with women who gave birth following fertility treatment (0.11 v. 0.08 events per 100 patient-years; adjusted HR 1.42, 95% CI 1.06–1.92) (Figure 1). A check of the consistency of the HR over time with the introduction of a time-dependent interaction term was significant (p = 0.002), which suggested departure from the proportional hazards assumption. In particular, the first year of follow-up showed the largest relative risk associated with fertility therapy failure for the composite of hospital admission for a cardiovascular event or death (adjusted HR 4.24, 95% CI 1.90–9.44). The trajectory thereafter of cardiovascular risk following fertility therapy failure ap-

### Table 1: Baseline characteristics of 28 442 women who received fertility therapy*

| Characteristic                                      | Failure n = 19 093 | Success n = 9349 | p value |
|-----------------------------------------------------|--------------------|------------------|---------|
| Age, yr, median (IQR)                               | 36 (32–39)         | 34 (31–37)       | < 0.001 |
| No. of fertility cycles (IQR)                       | 3 (1–5)            | 3 (2–6)          | < 0.001 |
| In vitro fertilization procedure                    | 2211 (11.6)        | 968 (10.4)       | < 0.001 |
| Rural residence                                     | 787 (4.1)          | 400 (4.3)        | < 0.001 |
| Income quintile                                     |                    |                  | < 0.001 |
| 1 (lowest)                                          | 3202 (16.8)        | 1048 (11.2)      |         |
| 2                                                   | 3477 (18.2)        | 1470 (15.7)      |         |
| 3                                                   | 3718 (19.5)        | 1877 (20.1)      |         |
| 4                                                   | 4221 (22.1)        | 2354 (25.2)      |         |
| 5 (highest)                                         | 4416 (23.1)        | 2591 (27.7)      |         |
| Unknown                                             | 59 (0.3)           | 9 (0.1)          |         |
| Prior obstetric delivery                            | 3167 (16.6)        | 1700 (18.2)      | < 0.001 |
| Any visit to emergency department in prior year     | 903 (4.7)          | 688 (7.4)        | < 0.001 |
| Any hospital admission in prior year                | 4193 (22.0)        | 2277 (24.4)      | < 0.001 |
| History of miscarriage                             | 3095 (16.2)        | 1700 (18.2)      | < 0.001 |
| History of stillbirth                               | 50 (0.3)           | 63 (0.7)         | < 0.001 |
| Hypertension, including gestational hypertension    | 729 (3.8)          | 656 (7.0)        | < 0.001 |
| Diabetes, including gestational diabetes             | 599 (3.1)          | 964 (10.3)       | < 0.001 |
| Hypercholesterolemia                                | 522 (2.7)          | 98 (1.0)         | < 0.001 |
| Smoking                                             | 34 (0.2)           | 6 (0.1)          | 0.02    |
| Obesity                                             | 435 (2.3)          | 125 (1.3)        | < 0.001 |
| Asthma                                              | 792 (4.1)          | 334 (3.6)        | 0.02    |
| Neoplasm                                            | 454 (2.4)          | 90 (1.0)         | < 0.001 |
| Depression                                          | 585 (3.1)          | 244 (2.6)        | 0.03    |
| Systemic autoimmune rheumatic disease               | 843 (4.4)          | 367 (3.9)        | 0.3     |
| Atrial fibrillation                                 | 267 (1.4)          | 165 (1.8)        | 0.02    |
| Ischemic heart disease                              | 174 (0.9)          | 62 (0.7)         | 0.03    |
| Cardiomyopathy                                      | 21 (0.1)           | 32 (0.3)         | < 0.001 |
| Cerebrovascular disease                             | 50 (0.3)           | 27 (0.3)         | 0.7     |
| Thromboembolic disease                              | 10 (0.1)           | 9 (0.1)          | 0.2     |
| Pulmonary circulation disease                       | 25 (0.1)           | 13 (0.1)         | 0.9     |

Note: IQR = interquartile range.

*Baseline characteristics were ascertained during the 2 years before the date of final fertility treatment except for history of live births, miscarriages and stillbirths, data for which were ascertained over the duration of available data.

†Unless stated otherwise.
approached that of cardiovascular risk following successful fertility therapy by about 5 years after final treatment. Restriction of the survival analysis to focus only on the time to first hospital admission for a cardiovascular event resulted in similar results, with the highest relative risk in the first year of follow-up (adjusted HR 3.21, 95% CI 1.24–8.34).

We observed no statistically significant heterogeneity in cardiovascular risk associated with fertility therapy failure across subgroups (all interaction p values > 0.1), except possibly according to history of miscarriage (interaction p = 0.06). In this subgroup, the increased risk appeared accentuated among women with a history of miscarriage (Figure 2). Of note, no subgroup showed a lower subsequent risk of adverse cardiovascular events associated with fertility therapy failure, including women who underwent only a single cycle of treatment (Figure 3).

Interpretation

In our cohort of more than 28 000 women, fertility therapy failure was common and was associated with an increased risk of adverse cardiovascular events. The largest relative increases were for heart failure and cerebrovascular events. The overall higher relative risk of cardiovascular events associated with fertility therapy failure was observed in diverse demographic subgroups, irrespective of the number of fertility treatment cycles and regardless of whether the diagnosis was primary or secondary infertility. The increased risk was particularly high among women with a prior miscarriage. The absolute risk was modest and the relative risk was mostly confined to the first 5 years following fertility therapy.

Several mechanisms may underlie our findings, as suggested by past studies linking fertility therapy, ovarian hyperstimulation and vascular complications.10,11 Prior population-based reports of women receiving fertility therapy (including in vitro fertilization) indicated an increased risk of pulmonary embolism throughout pregnancy, post partum10,11 and over subsequent years.12 An increased risk of thrombosis persists after the birth of a child conceived with the help of fertility therapy and may extend to women following fertility therapy failure. The risk may become increasingly relevant as more women with comorbidities contemplate reproduction with fertility therapy.35,36 The risk may also become more relevant as specific employers propose oocyte cryopreservation for future fertility as a workplace benefit.37,38
The cause of cerebrovascular events and heart failure following fertility therapy failure is likely complex.\textsuperscript{8,9,39} One potential mechanism is direct activation of the renin–angiotensin system with ovarian hyperstimulation,\textsuperscript{13–15} which influences systemic vascular permeability, sodium balance and blood pressure.\textsuperscript{16} Fluid shifts may result in depletion of intravascular volume and interstitial edema that mediates ischemic stroke, heart failure and venothromboembolic risks during pregnancy and potentially thereafter.

Whether these cardiovascular events could be prevented by long-term renin–angiotensin inhibitor therapy is a topic for future research, although some short-term success has been described for the early adverse effects of ovarian hyperstimulation.\textsuperscript{40,41} Additional potential options to protect against thromboembolism might include the control of risk factors or prophylactic antiplatelet therapy.\textsuperscript{42} In contrast, an observed decreased risk of coronary events following fertility therapy failure appears inconsistent with the hypothesis that endothelial dysfunction may already be present in women at risk of miscarriage and may become accentuated by fertility therapy failure.\textsuperscript{19}

Alternatively, fertility treatment may be innocuous and merely unmask a latent predisposition to premature cardiovascular disease among individuals at risk of infertility.\textsuperscript{22–25,43} Our prior research showed that women who delivered following fertility therapy had more short-term adverse cardiometabolic events but lower long-term cardiovascular risk compared with women who

![Figure 2: Risk of cardiovascular events or death following fertility therapy failure versus fertility therapy success, stratified by baseline risk factors. Adjusted rate ratios greater than 1.0 indicate an increased risk following fertility therapy failure. Error bars = 95% confidence intervals, CI = confidence interval.](image)

![Figure 3: Cumulative incidence of major cardiovascular events or death per 1000 patient-years following fertility therapy failure or success, stratified by number of treatment cycles.](image)
gave birth without assisted reproductive treatment. In the current study, we reduced confounding by the propensity of healthier women to receive fertility therapy, a bias that might mask potential detrimental biologic consequences. Together, these findings are consistent with the hypothesis that fertility therapy failure may represent an early indication for future cardiovascular disease because it represents a unique cardiometabolic stress test. As a result, the time following fertility therapy may be an opportunity to identify young women at increased risk of future cardiovascular events and initiate early strategies to modify risk factors (an opportunity that is currently unrecognized). This interpretation, however, warrants confirmation in further studies.

Limitations
Our approach provided a comprehensive evaluation of women following fertility therapy, yet the results should be interpreted with caution. In particular, we could not ascertain less invasive forms of fertility therapy, and information was not available about other clinical factors not detailed in administrative data, such as blood pressure, cholesterol levels and ventricular function. We could not distinguish among early types of fertility therapy failure and a small number of women who may have undergone ovarian induction for fertility preservation or egg donation. Although we could not definitively address whether successive cycles of fertility therapy were associated with increased cardiovascular risk, future studies using propensity-based methodology seem warranted to minimize selection bias and reduce confounding by indication. Finally, we combined diverse cardiovascular events for our composite outcome, which should be interpreted with caution because underlying mechanisms and risk factors may be different.

Conclusion
Fertility therapy failure was associated with an increased risk of long-term adverse cardiovascular events. A potential increase in cardiovascular events may become increasingly relevant with broader utilization of fertility therapy and longer follow-up. More informed decision-making around reproductive technology requires an awareness of potential risks and the need for continued long-term clinical care. Fertility therapy failure may be another sex-specific cardiovascular risk factor that warrants further research to identify women who may benefit from a cardiovascular risk evaluation.

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