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Mortality in a large community-based cohort of inner-city residents in Vancouver, Canada

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Abstract

Background: The Downtown Eastside is a robust and densely populated neighbourhood in Vancouver, Canada, that is characterized by low-income housing and drug use and a high prevalence of HIV infection. We evaluated mortality and excess mortality among the broader community of individuals living in this neighbourhood.

Methods: The Community Health and Safety Evaluation is a community-based study of inner-city residents in the Downtown Eastside who were recruited in 2003 and 2004. Participants' data were linked with data in provincial virology and mortality databases retrospectively and prospectively for the period 1991–2009. Mortality and standardized mortality ratios (SMRs) were calculated for the period 2003–2009 to compare death rates in the study population with rates in the population of Vancouver.

Results: Among 2913 participants, 374 deaths occurred, for an all-cause mortality of 223 per 10 000 person-years (95% confidence interval [CI] 201–247 per 10 000 person-years). Compared with the population of Vancouver, significant excess mortality was observed in the study population (SMR 7.1, 95% CI 6.4–7.9). Excess mortality was higher among women (SMR 15.4, 95% CI 12.8–18.5) than among men (SMR 5.8, 95% CI 5.1–6.6). Although crude mortality increased with age, excess mortality was greatest among participants less than 35 years old (SMR 13.2, 95% CI 9.4–18.5) and those 35–39 years old (SMR 13.3, 95% CI 10.3–17.1). Excess risk was also elevated among participants with hepatitis C virus (HCV), HIV and HCV/HIV infection, with SMRs of 5.9 (95% CI 4.9–7.1), 19.2 (95% CI 12.8–28.9) and 23.0 (95% CI 19.3–27.4), respectively.

Interpretation: Our study showed high mortality in this inner-city population, particularly when compared with the general population of Vancouver. Excess mortality was highest among women, younger participants and those infected with either HCV or HIV or both.

he Downtown Eastside is an inner-city neighbourhood in Vancouver, Canada, with a population of more than 16 000, including an estimated 4700 who are injection drug users.^{1,2} Rising rates of fatal drug overdose and HIV infection in this neighbourhood prompted the declaration of a public health emergency in 1997.^{3,4} Despite intensive study, improvements in social services and the implementation of several effective public health interventions,^{5,6} the impact on health outcomes and overall mortality at a community level is uncertain. Surveillance of mortality in inner-city populations is crucial to document disease trends and implement interventions that can have the biggest health impact.

Most studies of mortality in the Downtown Eastside have examined cohorts of injection drug users,^{2,7-9} which may not be generalizable to the broader community. Studies examining excess mortality have largely focused on all-cause mortality in specific populations, such as women⁷ and young injection drug users,² or have included surrounding neighbourhoods with higher socioeconomic status.¹⁰ We conducted this study to evaluate the mortality and excess mortality for all-cause and cause-specific deaths during the period 2003–2009 among residents in this inner-city neighbourhood.

Methods

Study population

The Community Health and Safety Evaluation (CHASE) cohort was designed to evaluate the uptake of health services and health outcomes in Vancouver's Downtown Eastside. To collect a broad sample of residents in this community, facility-based sampling¹¹ was used and venues for recruitment

Competing interests: See end of article.

This article has been peer reviewed.

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CMAJ Open 2013.DOI:10.9778/cmajo.20130002

were selected based on census tract data.¹² People were informed of the project through community-based agency staff, postings in local agencies, door-to-door initiatives and word of mouth. Surveys were administered in a variety of settings, including 10 community-based agencies, 2 community health clinics, 117 single-room occupancy hotels and social housing buildings, and a large space that operates as a needleexchange site. All residents included in the study had their names and personal health numbers verified through the British Columbia Ministry of Health database to facilitate linkage with hepatitis C virus and HIV test results and with health indicator databases.

Between January 2003 and December 2004, a total of 2913 participants received \$10 to complete a one-time, intervieweradministered survey (to collect information on demographics, health services utilization and drug use in the 6 months before the survey) and consented to have their data linked to health databases by name, date of birth and provincial health number. This study was approved by the University of British Columbia Providence Health Care Research Ethics Board.

Data collection

Participants' data were linked with data in provincial virology and mortality databases retrospectively and prospectively for the period 1991–2009. Linkage was performed using the participants' provincial health number or, for those with no match, their name and date of birth.¹² Results of antibody testing for hepatitis C virus (HCV) and HIV from 1991 to 2009 were obtained through database linkages with the British Columbia Centre for Disease Control and the University of British Columbia Virology Department (responsible for all HCV and HIV testing in the province).

For deaths from 2003 to 2009, the underlying causes of death were obtained from the BC Vital Statistics database, which captures information on all deaths in the province. Causes of death were classified according to the International Statistical Classification of Diseases and Related Health Problems, 10th revision (ICD-10) codes. Causes of death were grouped according to ICD-10 chapter headings as well as by drug-related, HIV-related and liver-related causes. Drug-related deaths comprised mental and behavioural disorders due to psychoactive substance use (ICD-10 codes F11-16 and F19), accidental poisoning by drugs (X40-44), suicide by drugs (X60-64), assault by drugs and medicaments (X85), poisoning by drugs or medicaments undetermined if accidental or intentional (Y10-14), and adverse effects of drugs and medicaments (Y40-574, Y577-79, Y598 and Y880). HIV-related deaths were defined as any death with an HIVrelated ICD-10 code (B20-24), whereas liver-related deaths comprised deaths from viral hepatitis (B15-19), liver cancer (C22), alcoholic liver disease (K70) or nonalcoholic liver disease (K71-77).

Outcome measures

The primary outcome measure was mortality. We determined both overall and cause-specific mortality and expressed them as rates of death and standardized mortality ratios (SMRs).

Statistical analysis

Participants who had linked data for HCV and HIV test results were divided into 4 mutually exclusive strata (HCV positive/HIV negative; HIV positive/HCV negative; HCV and HIV positive; and HCV and HIV negative) for stratified analyses. The HCV positive/HIV negative group included individuals with positive HCV test results and negative or unknown HIV results. The HIV positive/HCV negative group included those with positive HIV test results and negative or unknown HCV results. Participants with unknown test results were included in these groups because they had characteristics similar to those who tested negative (unpublished data). Causes of death were examined overall and among these

Table 1: Characteristics of 2913 participants in a communitybased cohort in the inner city of Vancouver

Characteristic	No. (%) of participants* n = 2913
Age, yr, mean ± SD	42.5 ± 10.1
Female sex	845 (29.0)
Aboriginal ethnicity	895 (30.7)
Unstable housing†	1801 (61.8)
Jail time†	610 (20.9)
Alcohol use†	1490 (51.2)
Drug use (injection or noninjection)†	2337 (80.2)
Injection drug use†	1114 (38.2)
Overdose†	205 (7.0)
Methadone treatment†	561 (19.3)
HCV infection‡	
Yes	1533 (52.6)
No	872 (29.9)
Unknown	508 (17.4)
HIV infection‡	
Yes	535 (18.4)
No	1735 (59.6)
Unknown	643 (22.1)
HCV and HIV infection strata‡	
HCV+/HIV– §	1068 (36.7)
HCV– /HIV+¶	70 (2.4)
HCV+/HIV+	465 (16.0)
HCV-/HIV-	693 (23.8)
Unknown**	617 (21.2)

Note: HCV = hepatitis C virus, SD = standard deviation.

*Unless stated otherwise.

†In the 6 months before the survey (self-reported). ‡Based on all linked laboratory test results (1991–2009). Characteristics of

participants by HCV/HIV infection strata are shown in Appendix 1 (available at www.cmajopen.ca/content/1/2/E68/suppl/DC1).

§Includes participants whose HCV result was positive and whose HIV status was either negative or unknown.

Includes participants whose HIV result was positive and whose HCV status was either negative or unknown.

**Includes participants whose HCV status and HIV status were both unknown; it also includes participants whose status was unknown for either HCV or HIV and who had a negative HIV or HCV result, respectively.

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strata. We calculated mortality using person-time methods. Person-years at risk were calculated for each participant as the time from the date of the survey questionnaire to the date of death or Dec. 31, 2009, whichever came first.

We calculated SMRs to compare death rates in the study population with rates in the population of Vancouver (specifically, the Health Service Delivery Area of Vancouver, population 590 240) for each cause of death. The SMRs were adjusted for sex, 5-year age group and calendar year of death, with the last 2 variables treated as time-dependent covariates. Participants with missing data on date of birth (0.2%, n = 7) were excluded from SMR analyses. We used the Poisson exact method to calculate 95% confidence intervals (CIs) for incidence rates and SMRs. Trends in rates from 2003 to 2009 were examined using Poisson regression. We considered p values of less than 0.05 (2-sided test) to be statistically significant. Statistical analyses were performed using R (version 2.15; R Foundation for Statistical Computing, Vienna, Austria) and STATA (version 10.1; StataCorp, College Station, Texas).

Results

Characteristics of the 2913 participants recruited into the CHASE cohort in 2003–2004 are shown in Table 1. The mean age was 42.5 years, and 845 (29.0%) were female. Among the participants for whom results of antibody testing for HCV and HIV were available during the linkage period (1991–2009), 63.7% (1533/2405) had HCV infection, 23.6% (535/2270) had HIV infection, and 21.8% (465/2130) had both. Injection or noninjection drug use in the 6 months before the survey was reported by 2337 (80.2%) of the participants.

Between January 2003 and December 2009, 374 deaths were reported over 16 778 person-years of observation, for an all-cause mortality of 223 per 10 000 person-years (95% CI

201–247 per 10 000 person-years) (Table 2). According to HCV and HIV status, the rate of death per 10 000 personyears was 167 (95% CI 138–203) among participants with HCV infection alone, 641 (95% CI 426–964) among those with HIV alone, 515 (95% CI 432–613) among those with HCV/HIV co-infection and 107 (95% CI 79–143) among those with neither HCV nor HIV infection (Table 3). Compared with the population of Vancouver, significant excess mortality was observed in the study population (SMR 7.1, 95% CI 6.4–7.9) (Table 2). Significantly higher excess risk was observed among participants with HCV infection (SMR 5.9, 95% CI 4.9–7.1), HIV infection (SMR 19.2, 95% CI 12.8–28.9) and HCV/HIV co-infection (SMR 23.0, 95% CI 19.3–27.4) compared with those who had neither HCV nor HIV infection (SMR 3.5, 95% CI 2.6–4.7) (Table 3).

Causes of death for the whole study population and for subgroups stratified by HCV and HIV status are shown in Tables 2 and 3, respectively (corresponding data for participants classified as having "unknown" status are shown in Appendix 2, available at www.cmajopen.ca/content/1/2 /E68/suppl/DC1). Overall, of the 374 deaths, the most common causes were infection (25.9%), external causes (16.6%), respiratory disease (11.8%) and neoplasm (11.2%); external causes included accidents, assault and intentional self-harm. Infection-related deaths were predominantly related to HIV/AIDS (80.4%). Excess mortality related to respiratory disease was due primarily to pneumonia (48.9% [22/45]) and chronic lower respiratory diseases (31.1% [14/45]). The highest cause-specific SMRs overall were for "other" causes, infection, respiratory disease, and mental and behavioural disorders; the same 4 causes accounted for much of the greatest excess risk among participants with HCV, HIV and HCV/HIV infection as well. Causes of deaths classified as "other" included the ICD-10 code for "other ill-defined and

Table 2: Mortality and exces	ss mortality among the 2	2913 participar	ts, by cause of death	
Cause of death	ICD-10 codes	No. of deaths	Rate per 10 000 person-years (95% CI)	SMR* (95% CI)
All		374	223 (201–247)	7.1 (6.4–7.9)
Infection	A00–B99	97	58 (47–71)	20.2 (16.5–24.6)
Neoplasm	C00–D48	42	25 (18–34)	2.7 (2.0–3.7)
Endocrine	E00–90	4	2 (1–6)	2.4 (0.9–6.3)
Mental or behavioural	F00–99	13	8 (4–13)	11.1 (6.4–19.0)
Nervous system	G00–99	8	5 (2–10)	7.0 (3.5–14.0)
Circulatory system	100–99	41	24 (18–33)	4.1 (3.0–5.5)
Respiratory system	J00–99	44	26 (20–35)	12.1 (9.0–16.3)
Digestive system	K00–93	20	12 (8–18)	7.3 (4.7–11.3)
Genitourinary system	N00–99	4	2 (1–6)	7.3 (2.7–19.4)
Other†	R00–99	39	23 (17–32)	37.8 (27.6–51.7)
External cause	V00-Y98	62	37 (39–47)	6.5 (5.0–8.3)

Note: CI = confidence interval; ICD-10 = International Statistical Classification of Diseases and Related Health Problems, 10th revision; SMR = standardized mortality ratio.

*Reference group was the population of Vancouver tincludes unknown causes of death

Table 3: Mort	ality an	oxe pr	cess mor	rtality by ca	iuse of d	leath in	subgro	ups stra	tified by HCV and	HIV sta	itus*					
			HCV+/H <i>n</i> = 10	IV-† 168			-	HCV-/HIV <i>n</i> = 70	++			HCV+/HI <i>n</i> = 465	+		HCV–/H <i>n</i> = 69	IV- 3
Cause of death	No. of deaths	- 6)	Rate§ 5% CI)	SMF (95%	ر ت	No. of deaths	(95 [°]	ate§ % CI)	SMR (95% CI)	No. of deaths	н <u>(</u>)	łate§ 5% CI)	SMR (95% CI)	No. of deaths	Rate§ (95% CI)	SMR (95% CI)
AII	106	167 ((138–203)) 5.9 (4.9	-7.1)	23	641 (42	26–964)	19.2 (12.8–28.9)	126	515 (4	32-613)	23.0 (19.3–27.4)	44	107 (79–143)	3.5 (2.6–4.7)
Infection	10	16	(8–29)	5.4 (2.9	-10.1)	13	362 (2	10-624)	130.3 (75.7-224.5)	68	278 (2	19-352)	107.7 (84.9–136.6)	4	10 (4–26)	3.5 (1.3–9.4)
Neoplasm	6	14	(7–27)	1.8 (0.9	-3.4)	0		1	I	9	25 ((11–55)	4.1 (1.9–9.2)	:	27 (15–48)	2.9 (1.6–5.3)
Endocrine	0		I	I		N	.) 95	14–223)	50.0 (12.5-200.0)	0		I	I	0	I	I
Mental and behavioural	4	9	(2–17)	10.2 (3.8	-27.2)	-	28	(4–200)	37.7 (5.3–267.5)	9	25 ((11–55)	50.5 (22.7–112.4)	-	2 (0–17)	3.7 (0.5–26.6)
Nervous system	0	ε	(1-13)	5.4 (1.4	-21.6)	0		1	I	2	ω	(2–33)	18.8 (4.7–75.3)	с	7 (2–23)	11.5 (3.7–35.5)
Circulatory system	14	22	(13–37)	4.4 (2.6	-7.5)	-	28	(4–198)	4.1 (0.6–28.8)	ę	12	(4–38)	3.5 (1.1–10.9)	7	17 (8–36)	2.9 (1.4–6.1)
Respiratory system	19	30	(19–47)	16.4 (10.4	-25.7)	-	28	(4–198)	11.6 (1.6–82.2)	œ	33 ((16–65)	24.5 (12.3–49.0)	വ	12 (5–29)	6.0 (2.5–14.3)
Digestive system	œ	13	(6–25)	8.4 (4.2	-16.7)	0		I	I	ო	12	(4–38)	10.4 (3.4–32.2)	-	2 (0–17)	1.5 (0.2–10.6)
Genitourinary	0		I	I		0		I	I	0		I	I	-	2 (0–17)	8.4 (1.2–59.4)
Other	12	19	(11–33)	30.4 (17.3	-53.6)	-	28	(4–198)	46.9 (6.6–333.1)	12	49	(28-86)	89.9 (51.1–158.3)	9	15 (7–32)	23.8 (10.7-53.1)
External	28	44	(31–64)	7.6 (5.2	-11.0)	4	111 (42–297)	21.2 (8.0–56.5)	18	74 ((46–117)	13.9 (8.8–22.1)	വ	12 (5–29)	2.2 (0.9–5.2)
Note: CI = confid. Causes of death HCV result, resper Thncludes particip Andre particip SRate per 10 000	ence inter for partic ictively) a ants who ants who person-y	rval, HC cipants are sho se HC se HIV /ears. s of dee	ZV = hepati in the "unkı wn in Appe V result was result was	itis C virus, SN nown status" g ndix 2 (availab s positive and v positive and w	IR = standa roup (i.e., tl le at www.c whose HIV /hose HCV	ardized mo hose whos cmajopen status wa status wa	rtality rai se HCV s ca/conte s either r s either r	tio. status and H nt/1/2/E68/s negative or u negative or u	IIV status were both un suppl/DC1). Jnknown. Jnknown.	known, ar	nd those	whose statu	s was unknown for eith	er HCV ol	r HIV and who had	d a negative HIV or

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unspecified causes of mortality" (n = 34) and deaths with missing data regarding cause of death (n = 6).

Mortality and excess mortality related to drug use, liver disease and HIV infection are shown in Tables 4 and 5 and in Appendix 3 (available at www.cmajopen.ca/content/1/2/E68/suppl/DC1). Overall, 14.2% of deaths were drug-related (32 deaths [95% CI 24–41] per 10 000 person-years), which represents a considerable excess risk of drug-related mortality (SMR 12.0, 95% CI 9.1–15.7). Liver-related deaths accounted for 5.9% of the deaths, for a rate of 13 (95% CI 9–20) per 10 000 person-years (SMR 5.9, 95% CI 3.9–8.9). The rate of HIV-related deaths was 46 (95% CI 37–58) per 10 000 person-years (SMR 24.9, 95% CI 20.0–31.1).

Women in the cohort had rates of death similar to those among men (Table 4). However, their excess mortality was significantly higher than the excess mortality among men for all causes (SMR 15.4 [95% CI 12.8–18.5] v. 5.8 [95% CI 5.1–6.6]) and for drug-, liver- and HIV-related deaths (Table 4).

Although crude mortality increased with age, excess mortality was highest among participants less than 35 years old (SMR 13.2, 95% CI 9.4–18.5) and among those 35–39 years old (SMR 13.3, 95% CI 10.3–17.1); the corresponding SMRs and 95% CIs for participants aged 45–49 years and those 50 years and older were 6.0 (4.7–7.6) and 5.3 (4.5–6.3) (Appendix 4, available at www.cmajopen.ca/content/1/2/E68/suppl/DC1). Liver-related mortality also increased with age, but it was associated with a relatively stable excess risk across age groups (Figure 1). In contrast, drug-related mortality and excess mortality

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tended to peak at age 35–39 years. HIV-related mortality was greatest among those aged 45–49 years, with a secondary peak at age 35–39, but HIV-related SMRs trended downward overall with age. The annual rates of death over the period 2003–2009 did not show a significant linear trend (Figure 2; p = 0.6).

Interpretation

This large study of a cohort of inner-city residents in Vancouver, Canada, demonstrated several important findings. Very high rates of death, particularly when compared with rates for the general population of Vancouver, were evident for all-cause and cause-specific mortality. Excess mortality was highest among women, younger participants and participants infected with HCV or HIV or both. Major causes of death included HIV infection, drug-related deaths and unknown causes.

The all-cause mortality was high (223 per 10 000 personyears) and was associated with considerable excess mortality compared with the general Vancouver population (SMR 7.1). This ratio is much greater than the SMR of 1.2 reported by the British Columbia Vital Statistics Agency for the local health area that contains the Downtown Eastside;¹⁰ the difference is likely because the latter includes wealthier neighbourhoods outside the inner city that decrease the observed rates of death. The SMR in the cohort inner-city residents in our study is closer in magnitude to those in populations exclusively composed of injection drug users. Previous studies of mortality among injection drug users reported SMRs of

HIV INTECTION among	j the 2913 participants		
Cause of death	No. of deaths	Rate per 10 000 person-years (95% CI)	SMR† (95% CI)
All	374	223 (201–247)	7.1 (6.4–7.9)
Men	263	220 (195–248)	5.8 (5.1–6.6)
Women	111	230 (191–277)	15.4 (12.8–18.5)
Drug-related‡	53	32 (24–41)	12.0 (9.1–15.7)
Men	36	30 (22–42)	9.3 (6.7–12.9)
Women	17	35 (22–57)	30.3 (18.8–48.7)
Liver-related§	22	13 (9–20)	5.9 (3.9–8.9)
Men	15	13 (8–21)	4.5 (2.7–7.4)
Women	7	15 (7–30)	17.9 (8.5–37.6)
HIV-related¶	78	46 (37–58)	24.9 (20.0–31.1)
Men	54	45 (35–59)	18.7 (14.4–24.5)
Women	24	50 (33–74)	96.8 (64.9–144.5)

Table 4: Mortality and excess mortality related to drug use, viral hepatitis and liver disease, and

Note: CI = confidence interval; ICD-10 = International Statistical Classification of Diseases and Related Health Problems, 10th revision; SMR = standardized mortality ratio.

*Causes of death for participants in the "unknown status" group (i.e., those whose HCV status and HIV status were both unknown, and those whose status was unknown for either HCV or HIV and who had a negative HIV or HCV result, respectively) are shown in Appendix 3 (available at www.cmajopen.ca/content/1/2/E68/suppl/DC1).

†Reference group was the population of Vancouver.

[‡]Deaths from mental and behavioural disorders due to psychoactive substance use (ICD-10 codes F11–16, F19), accidental poisoning by drugs (X40–44), suicide by drugs (X60–64), assault by drugs and medicaments (X85), poisoning by drugs or medicaments undetermined if accidental or intentional (Y10–14), and adverse effects of drugs and medicaments (Y40–574, Y577–79, Y598, Y880).

Speaths from viral hepatitis (B15–19), liver cancer (C22), alcoholic liver disease (K70) and nonalcoholic liver disease (K71–77).

¶ICD-10 codes B20-24.

Table 5: Mo	rtality a	ind excess mo	ortality related to	drug u	se, viral hepati	itis and	liver disease,	, and HI	/ infection in su	ıbgroups stratified b	y HCV ar	nd HIV status*	
		HCV+/HI ⁿ n = 106	V-† 38		HCV-/F	+1/V+# 70			HCV+/F n = 4	11V+ 35		HCV-/HIV. <i>n</i> = 693	1
Cause of death	No. of deaths	Rate§ (95% CI)	SMR (95% CI)	No. of deaths	Rate§ (95% CI)	5)	SMR 95% CI)	No. of deaths	Rate§ (95% CI)	SMR (95% CI)	No. of deaths	Rate§ (95% CI)	SMR (95% CI)
AII	106	167 (138–203)	5.9 (4.9–7.1)	23	641 (426–964)	19.2	(12.8–28.9)	126	515 (432–613)	23.0 (19.3–27.4)	44	107 (79–144)	3.5 (2.6–4.7)
Men	75	169 (135–233)	4.8 (3.8–6.0)	18	794 (500–1261)	17.1	(10.8–27.2)	17	501 (401–627)	16.9 (13.5–21.1)	34	112 (80–156)	3.0 (2.2–4.2)
Women	31	164 (115–233)	13.1 (9.2–18.6)	5	378 (157–908)	34.4	(14.3-82.7)	49	538 (407–712)	53.8 (40.6–71.2)	10	93 (50–173)	7.6 (4.1–14.2)
Drug-related	22	35 (23–53)	12.6 (8.3–19.1)	0	56 (14–223)	23.1	(5.8–92.3)	21	86 (56–132)	33.7 (22.0–51.7)	4	10 (4–26)	3.8 (1.4–10.2)
Men	16	36 (22–59)	10.5 (6.4–17.1)	2	88 (22–535)	27.8	(7.0-111.3)	÷	72 (40–129)	21.2 (11.7–38.2)	4	13 (5–35)	4.3 (1.6–11.5)
Women	9	32 (14–71)	26.9 (12.1–59.9)	0	I		I	10	110 (59–204)	96.8 (52.1–180.0)	0	I	I
Liver-related**	13	21 (12–35)	9.6 (5.6–16.5)	0	1		I	5	20 (9–49)	12.1 (5.0–29.1)	0	I	I
Men	6	20 (11–39)	7.4 (3.8–14.2)	0	I		I	0	13 (3–52)	5.6 (1.4–22.3)	0	I	I
Women	4	21 (8–56)	28.2 (10.6–75.1)	0	I		I	e	33 (11–102)	54.4 (17.6–168.8)	0	I	I
HIV-related††	0	1	1	13	362 (210-624)	199.8 (116.0–344.1)	64	262 (205–334)	142.8 (111.8–182.5)	0	I	I
Men	0	1	1	10	441 (237-820)	177.0	(95.2–329.0)	43	280 (208–377)	108.0 (80.1–145.6)	0	I	I
Women	0	I	I	e	227 (73–703)	349.9 (112.8–1100.0)	21	231 (150–354)	420.7 (274.3–645.2)	0	I	I
Note: CI = confi Causes of dea HCV result, res HICUdes partic #Includes partic §Rate per 10 00 ¶Deaths from m **Deaths from v 1†1CD-10 code	idence int idence int pectively) zipants wh zi parts wh zi parta and g by drugs <i>i</i> ral hepat. s B20-24.	erval; ICD-10 = Int licipants in the "unh are shown in Appu ose HCV result wa vears. Vears. I behavioural disor i behavioural disor i s or medicaments i its (B15-19), liver	ternational Statistical (known status" group (endix 3 (available at v as positive and whose s positive and whose orders due to psychoad undetermined if accidu cancer (C22), alcoho	Classifica i.e., those www.cmaj MLCV statt HCV statt tive subst antal or ir lic liver di	tion of Diseases an whose HCV status open.ca/content/1/, us was either negat us was either negat us was either negat ance use (ICD-10 tentional (Y10-14) sease (K70) and nc	d Related s and HIV 2/E68/supt tive or unk tive or unk tive or unk tive or adve and adve onalcoholic	Health Problems status were both p/DC1). nown. nown. -16, F19), accide srse effects of dru- srse effects of dru- srse effects of dru-	, 10th revisuunknown, unknown, ental poiso gs and me 71–77).	sion; SMR = standar and those whose st ing by drugs (X40- odicaments (Y40-57	dized mortality ratio. atus was unknown for eithe 44, suicide by drugs (X60- 4, Y577-79, Y598, Y880).	r HCV or HI -64), assaul	IV and who had a t by drugs and me	negative HIV or dicaments

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10-15,^{13,14} and excess mortality among users of opiates and opioid substitution therapy has been in the range of 5-13.¹⁵⁻¹⁷

Most of the causes of death associated with the greatest excess mortality, including infection, "other" causes and mental or behavioural disorders, were consistent with major causes of death identified in other studies.^{14,15,18-23} Of note was the elevated excess mortality related to respiratory disease (primarily pneumonia). A study involving drug users entering treatment in Rome, Italy, reported a similar SMR of 10.2 for respiratoryrelated death.²³ This is consistent with the observation that pneumonia is an important cause of morbidity and hospital admission among injection drug users.²⁴⁻²⁶

The very high drug-related mortality and excess mortality in our study are consistent with prior findings among injection drug users in Vancouver⁸ and elsewhere.^{13,15,23,27,28} Liver-



Figure 1: (A) Mortality and (B) excess mortality related to drug use, viral hepatitis and liver disease, and HIV infection among 2913 participants in a large community-based cohort in the inner city of Vancouver, 2003–2009, by age group. Drug-, liver- and HIV-related deaths are defined in Methods. The reference group for the standardized mortality ratios was the population of Vancouver.

related mortality, which accounted for 6% of the deaths, was associated with an SMR of 5.9. Comparatively higher SMRs were reported in studies involving people taking opioid substitution therapy (SMR 17.0)¹⁶ and among people infected with hepatitis B virus (SMR 10.0) and HCV (SMR 15.8).²⁹ Population differences such as age, proportion with viral hepatitis infection and pattern of alcohol use may account for such differences in risk. HIV-related mortality and excess mortality were very high in our study, particularly among participants known to have HIV infection, among whom 77 (51.7%) of 149 deaths were HIV-related. In studies involving injection drug users, deaths caused by HIV/AIDS accounted for 34%–38% of deaths among HIV-infected people.^{8,14}

Sex-related differences were evident in the all-cause mortality results: although men and women had similar rates of death, women had a significantly higher excess mortality (SMR 15.4 v. 5.8 for men). SMRs for drug-, liver- and HIVrelated deaths followed similar patterns. In a meta-analysis of mortality among opioid users, men had higher crude mortality rates than women, but women had higher SMRs.¹⁸ A similar pattern in the SMRs for all-cause mortality was observed in a cohort of Vancouver injection drug users.²⁷ The higher excess mortality among women likely reflects the fact that women in the general population tend to have a lower mortality than men for many major causes of death.

The increase in crude mortality from the youngest group to the oldest is consistent with expected increases in diseases of aging, including malignant diseases and cardiovascular diseases. The lower excess mortality among older participants could be explained by concurrent increases in these agerelated conditions in the general population and by survivorship bias. Liver-related mortality increased with age, which has been observed previously in the HCV-infected subset of this cohort.³⁰ Among younger inner-city residents, liverrelated deaths contributed minimally to mortality, whereas among people aged 50 years or older, the rate of liver-related deaths was similar to the drug- and HIV-related rates.



Figure 2: Mortality in the inner-city cohort, by year of follow-up. Drug-, liver- and HIV-related deaths are defined in Methods. Error bars = 95% confidence intervals.

Limitations

This study has several limitations. Because the marginalized, hard-to-reach population under study did not allow for randomized sampling, the sample cannot be considered representative,11 as in other studies with similar populations.31 However, the sampling measures used ensured a broad reach and inclusion of participants and provided the highest possible degree of representativeness under real-life circumstances. Participation bias related to illicit activities or status in the study population may also be an issue,¹¹ although this bias is also found in general population surveys.³² The study relied on participants' self-report for some data elements (e.g., history of drug use), the validity of which cannot be formally ascertained. However, self-reported data in similar studies have been shown to be sufficiently valid.33 This is an observational cohort study, and no conclusions should be drawn with regard to causality given the possibility of unmeasured confounding variables. Data regarding HCV and HIV infection in this cohort were derived from tests ordered for clinical indications; systematic testing was not performed. HCV status was unknown for 17% of the participants and HIV status for 22%; however, their demographic characteristics and rates of death were similar to those for participants who had neither HCV nor HIV infection. Participants with positive HCV test results but no linked HIV test results were considered negative for HIV, and those with positive HIV and unknown HCV were classified as HCV negative; although the baseline characteristics between the negative and unknown groups were similar in these cases, some HCV/HIV co-infected individuals may have been misclassified and mortality overestimated in the monoinfected groups. Linked HCV test results did not always include HCV RNA results, and spontaneous clearance may have occurred among some of those with presumed HCV infection, which led to an underestimation of the observed overall effect of HCV infection on mortality. Although the linked mortality records capture all deaths reported in the province of British Columbia, any deaths that went unreported or that occurred in other jurisdictions would lead to an overestimation of person-years of follow-up and an underestimation of mortality.

Conclusion

Our study showed high mortality in this inner-city population, particularly when compared with the general population of Vancouver. Excess mortality was highest among women, younger participants and those infected with HCV or HIV or both. The significant excess mortality and the leading causes of death observed in this study confirm the need for comprehensive, multidisciplinary services to support this high-risk population and optimize health outcomes, particularly among youth and women. Screening, follow-up and appropriate treatment for HIV and HCV infection are clearly important tools to reduce the risk of HIV-related mortality and transmission and to prevent HCV-related health sequelae. Measures such as mental health care, improved access to housing and harm reduction services are also important to address the drug-related deaths. Every interaction with the health care system can provide an opportunity to engage members of this inner-city population in the types of comprehensive care needed to address their diverse health conditions. Further investment in programs to improve inner-city health care (e.g., HIV, HCV, mental health and addiction services) may help to address the considerable morbidity and mortality in this inner-city population.

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Competing interests: Mel Krajden has received research grants, paid to his institution, from Merck, Roche, Siemens and Gen-Probe (Hologic). Gregory Dore is a member of advisory boards for Roche, Merck, Gilead, Bristol-Myers Squibb, AbbVie and Janssen; he has received research grants, paid to his institution, from Roche, Merck, Gilead, Bristol-Myers Squibb and AbbVie, and he has received payment for expert testimony from Janssen, speaker fees from Roche and Merck, and travel assistance from Roche, Merck and Gilead. Jason Grebely is a member of the advisory board for Merck; he has received a research grant, paid to his institution, from Merck. No competing interests declared by the other authors.

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Contributors: Jason Grebely and Mark Tyndall contributed equally to this work. Calvin Lai and Mark Tyndall designed the original study and wrote the protocol. Gregory Deans, Jesse Raffa, Jason Grebely and Mark Tyndall designed the mortality substudy. Gregory Deans, Jesse Raffa and Jason Grebely drafted the primary statistical analysis plan, which was reviewed by Calvin Lai, Benedikt Fischer, Mel Krajden, Janaki Amin, Scott Walter and Gregory Dore. The primary statistical analysis was conducted by Gregory Deans and was reviewed by all of the coauthors. All of the authors reviewed the data analysis. Gregory Deans wrote the first draft of the manuscript. All of the authors critically revised the manuscript and approved the final version submitted for publication.

Funding: The Community Health and Safety Evaluation cohort was established with funding from the Vancouver Coastal Health Authority. Gregory Deans was supported by funding from the National Canadian Research and Training Program in Hepatitis C. Gregory Dore is supported by a National Health and Medical Research Council Practitioner Research Fellowship. Jason Grebely is supported by a National Health and Medical Research Council Career Development Fellowship.

Supplemental information: For reviewer comments and the original submission of this manuscript, please see www.cmajopen.ca/content /1/2/E68/suppl/DC1