Recidivism and mortality after in-jail buprenorphine treatment for opioid use disorder

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ABSTRACT

Background: Buprenorphine is an effective medication for opioid use disorder (MOUD) when offered in community-based settings, but evidence is limited for incarcerated populations, particularly in relation to recidivism. In Massachusetts, Franklin County jail (FCSO) was among the first to provide buprenorphine; adjacent Hampshire County jail (HCHC) offered it more recently. These jails present a natural experiment to determine whether outcomes are different between individuals who did and did not have the opportunity to receive buprenorphine in jail.

Methods: We examined outcomes of all incarcerated adults with opioid use disorder (n = 469) who did (FCSO n = 197) and did not (HCHC n = 272) have the opportunity to receive buprenorphine. The primary outcome was post-release recidivism, defined as time from jail exit to a recidivism event (incarceration, probation violation, arraignment). Using Cox proportional hazards models, we investigated site as a predictor, controlling for covariates. We also examined post-release deaths.

Results: Fewer FCSO than HCHC individuals recidivated (48.2% vs. 62.5%; p = 0.003); fewer FCSO individuals were re-arraigned (36.0% vs. 47.1%; p = 0.046) or re-incarcerated (21.3% vs. 39.0%; p < 0.0001). Recidivism risk was lower in the FCSO group (hazard ratio 0.71, 95% confidence interval 0.56, 0.89; p = 0.003), net of covariates. We also examined post-release deaths.

Conclusions: Among incarcerated adults with opioid use disorder, risk of recidivism after jail exit is lower among those who were offered buprenorphine during incarceration. Findings support the growing movement in jails nationwide to offer buprenorphine and other agonist medications for opioid use disorder.

1. Introduction

Incarcerated individuals with opioid use disorder (OUD) are at high risk for overdose and other adverse outcomes after community release (Binswanger et al., 2013; Pizzicato et al., 2018). Medications to treat OUD (MOUD, i.e., buprenorphine, methadone, naltrexone) hold great promise to improve these outcomes among incarcerated populations (Mace et al., 2019; Malta et al., 2019; SAMHSA, 2019), but its implementation is not standard-of-care in U.S. jails and prisons (Grella et al., 2018; Macmadu et al., 2020; Simon et al., 2021). Most correctional facilities that do offer MOUD are large urban jails (e.g. New York City, San Francisco, Albuquerque) or part of unified state systems (e.g., Rhode Island, Vermont), and typically only offer naltrexone, with fewer facilities also offering buprenorphine (Wakeman and Rich, 2015). Prior studies on post-release outcomes have mostly examined the effects of methadone and naltrexone (Moore et al., 2019) and reported reduced overdoses, reduced risks for infectious disease, and other beneficial outcomes (Brinkley-Rubinstein et al., 2018; Fanucchi et al., 2019; Farrell-MacDonald et al., 2014; Haas et al., 2021; Murphy et al., 2017; Springer et al., 2018; Wilson et al., 2012). Few studies, however, have
examined the impact of buprenorphine in county jails on post-release recidivism (Moore et al., 2019), a crucial outcome to convince lawmakers and public safety officials of its value in correctional settings.

The Franklin County Sheriff’s Office (FCSO) in Greenfield, Massachusetts was among the first rural jails nationwide to offer buprenorphine (begun in February 2016), in addition to naltrexone, to incarcerated residents (Donelan et al., 2021). Hampshire County (located immediately south of Franklin County) has a similar rural jail, but the Hampshire County House of Corrections (HCHC) did not provide buprenorphine until May 2019. The current study uses the differences in buprenorphine provision between these jails in adjacent counties with similar recidivism rates as a natural experiment to examine the post-release recidivism and mortality outcomes of persons who were offered buprenorphine while incarcerated after their return to the community. We hypothesized that outcomes would be better among individuals who exited FCSO (i.e., offered buprenorphine pre-release) than among individuals who exited HCHC (i.e. not offered buprenorphine pre-release).

2. Methods

2.1. Study sample and data sources

The study sample includes all adults with opioid use disorder (OUD) who exited one of two participating jails between January 1, 2015 and April 30, 2019 (N = 469; 197 from FCSO, 272 from HCHC). Research staff extracted data from each jail’s electronic medical records (EMR) system to identify OUD diagnosis, receipt of MOUD while in jail, date of first jail exit (the “index jail episode,” for FCSO only we used first jail exit when MOUD was prescribed), and demographics. We verified OUD diagnosis and MOUD receipt by cross-checking EMR data against other records (prescription monitoring program; criminal justice records), and confirmed that no individuals in HCHC received MOUD while incarcerated (1 received naltrexone at release). In contrast, of individuals included in the FCSO group, 93.4% received MOUD while incarcerated, 1 was eligible for MOUD but did not receive it, and 12 could not be verified. Additionally, 53.1% of the FCSO group was inducted onto MOUD during incarceration, 38.8% continued MOUD at entrance per a prescription on file, and 8.2% had an unknown status. Most Franklin individuals received buprenorphine (86.2%), and fewer received extended-release naltrexone (7.1%), oral naltrexone (<1%), or an undocumented MOUD type (6.1%). Buprenorphine medication dosage ranged from 2 to 16 mg. For naltrexone, 1 person received 50 mg orally, documented MOUD type (6.1%). Buprenorphine medication dosage included in the FCSO group, 93.4% received MOUD while incarcerated, 1 was eligible for MOUD but did not receive it, and 12 could not be verified. Additionally, 53.1% of the FCSO group was inducted onto MOUD during incarceration, 38.8% continued MOUD at entrance per a prescription on file, and 8.2% had an unknown status. Most Franklin individuals received buprenorphine (86.2%), and fewer received extended-release naltrexone (7.1%), oral naltrexone (<1%), or an undocumented MOUD type (6.1%). Buprenorphine medication dosage ranged from 2 to 16 mg. For naltrexone, 1 person received 50 mg orally, and 14 received extended-release 380 mg intramuscularly.

To measure post-release outcomes, we obtained administrative data on the entire sample, ensuring that each individual had at least one year of observation after jail exit. Mean ±SD days from jail exit to end of observation was 618.8 ± 194.4 for the FCSO group (~20 months) and 745.2 ± 269.2 for the HCHC group (~25 months). We determined the one-year follow-up period for each individual based on the index release date.

The Baystate Health Institutional Review Board approved all study procedures and obtained federal certification for prisoner research.

2.2. Measures

2.2.1. Recidivism

Massachusetts Board of Probation (BOP) records contain statewide information on incarcerations, arraignments, and probation violations. Record reviews documented details such as type of event, date of occurrence, and number of events. We defined recidivism as any incarceration, arraignment, or probation violation occurring after release from the index jail episode.

2.2.2. All-Cause Mortality

Death information was searched on the web-based National Death Register, which provides the date of death, and by obtaining death certificates from state or county Vital Statistics offices for the cause of death.

2.3. Data analysis

We determined follow-up duration for our primary outcome using date of exit from the index jail episode to date of first recidivism event, death, or end of record review, whichever occurred first. Record review ended on April 30, 2020 to ensure all participants had at least 1 year of observation. Using a conservative, intent-to-treat approach, analyses treated individuals released from FCSO as having been offered MOUD, and vice-versa if released from HCHC. Cox proportional hazards (PH) models were fit to examine jail site as a predictor of recidivism after the index release, with covariates included based on baseline imbalances of demographic characteristics or criminal justice experiences. Number of prior incarcerations was highly correlated with number of prior arraignments and age of 1st arraignment. Thus, we selected as covariates number of prior incarcerations and index jail status is pre-trial vs. sentenced. Chi-square or t-tests compare characteristics at baseline by site. Hypothesis tests use a two-sided, significance level of α = 0.05.

We also compared the proportion of participants from each site who had any of the different types of recidivism events and arraignment charges using logistic regression models adjusted for baseline covariates. We limited these analyses to the first year after index jail release in order to 1. Investigate differences which may be most attributable to the MOUD intervention and 2. Eliminate bias due to the fact that HCHC participants, on average, had a longer record review time and would have more time to have recidivated.

We performed sensitivity analyses to explore the potential influence of within-site differences in site characteristics or durations of follow-up.

Table 1

<table>
<thead>
<tr>
<th>Demographics</th>
<th>Franklin County Sheriff’s Office (FCSO) N = 197</th>
<th>Hampshire County House of Corrections (HCHC) N = 272</th>
<th>P value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male – N (%)</td>
<td>179 (91.0)</td>
<td>272 (100)</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>Race – N (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>189 (96.0)</td>
<td>260 (96.0)</td>
<td>0.95</td>
</tr>
<tr>
<td>Black/AA</td>
<td>7 (4.0)</td>
<td>10 (4.0)</td>
<td></td>
</tr>
<tr>
<td>Other/Unknown</td>
<td>1 (1.0)</td>
<td>2 (1.0)</td>
<td></td>
</tr>
<tr>
<td>Age – Mean (sd)</td>
<td>34.5 (9.3)</td>
<td>35.1 (9.8)</td>
<td>0.49</td>
</tr>
<tr>
<td>Medication for Opioid Use Disorder (MOUD) – N (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Buprenorphine</td>
<td>170 (86.2)</td>
<td>0 (0.0)</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>Naltrexone</td>
<td>14 (7.1)</td>
<td>0 (0.0)</td>
<td></td>
</tr>
<tr>
<td>Undocumented MOUD type</td>
<td>12 (6.1)</td>
<td>0 (0.0)</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>1 (1.0)</td>
<td>272 (100)</td>
<td></td>
</tr>
<tr>
<td>CJS experiences before index jail episode</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First arraigned as a juvenile – N (%)</td>
<td>81 (41.0)</td>
<td>139 (51.0)</td>
<td>0.03</td>
</tr>
<tr>
<td>Age at first arraignment – Mean (sd)</td>
<td>19.2 (7.2)</td>
<td>17.7 (6.2)</td>
<td>0.02</td>
</tr>
<tr>
<td># of arraignments – Mean (sd)</td>
<td>11.3 (9.3)</td>
<td>15.7 (11.6)</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td># of incarcerations – Mean (sd)</td>
<td>3.0 (4.5)</td>
<td>4.9 (6.1)</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>CJS experiences on index jail episode</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jail status is sentenced – N (%)</td>
<td>38 (19.0)</td>
<td>113 (42.0)</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td># of days incarcerated – Mean (sd)</td>
<td>78.1 (126.1)</td>
<td>85.1 (131.1)</td>
<td>0.56</td>
</tr>
</tbody>
</table>

* t-test for continuous and chi square for categorical
of baseline differences between participants from FCSO and HCHC (see Table 1) by fitting a series of separate Cox PH models for time to 1st recidivism using restricted samples. We examined 3 sources of potential bias: 1. Gender, by excluding females; 2. Prior criminal justice system involvement, by excluding participants whose number of prior incarcerations were at or above the 75th percentile and those who were first incarcerated as a juvenile; and 3. Holding status at index jail stay, by excluding participants with sentenced status. In additional sensitivity analyses, we sought to confirm the effectiveness of buprenorphine by excluding FCSO participants who received naltrexone or whose MOUD type was unknown. Hazard ratios and confidence intervals were reported for each analysis.

3. RESULTS

3.1. Baseline characteristics

At baseline, demographics were similar across sites but participants differed in terms of interactions with the criminal justice system (Table 1). Most participants were male, White, and the mean age was 34.5 for FCSO and 35.1 for HCHC. Individuals in FCSO had fewer criminal justice interactions than those in HCHC, and these interactions began at an older age. Compared to individuals in HCHC, fewer individuals in FCSO were first arraigned as a juvenile and the FCSO group also had fewer prior arraignments and incarcerations. On the index jail episode, fewer individuals in FCSO than in HCHC had a sentenced status and more were pre-trial detainees.

3.2. Recidivism

Recidivism was defined as any incarceration, probation violation, or arraignment after index jail release (Table 2). Review and analysis of BOP records indicated that fewer individuals in FCSO than in HCHC recidivated (48.2% vs. 62.5%, respectively). Among people who did recidivate, the most common type of first recidivism event was an arraignment for both the FCSO and HCHC groups (67% vs. 71%), with fewer people recidivating with a re-incarceration (11% vs. 13%) or probation violation (22% vs. 16%) (data not shown).

The percentage of participants from FCSO experiencing arraignment or re-incarceration was approximately 11–18% lower than participants from HCHC. Individuals released from FCSO had reduced odds of any type of recidivism during the first year post-release (adjusted odds ratio 0.51 95% CI 0.35, 0.76; p = 0.001), and specifically, reduced odds of any post-release arraignment (aOR 0.67 95% CI 0.45, 0.99; p = 0.046) and incarceration (aOR 0.37 95% CI 0.24, 0.58; p < 0.0001). FCSO participants were less likely to have been arraigned on any property charges compared to HCHC participants (aOR 0.39 95% CI 0.22, 0.69; p = 0.001).

The mean±SD days from jail exit to first recidivism event was 132.9 ± 103.8 in FCSO and 129.4 ± 100.1 in HCHC. Cox proportional hazards model results showed a decreased risk of recidivism for FCSO compared to HCHC, with an unadjusted hazard ratio (95% confidence interval) of 0.71 (0.56, 0.89) and an adjusted HR (95% CI) of 0.68 (0.53, 0.86) (Fig. 1).

We conducted sensitivity analyses of recidivism hazard ratios (95% confidence interval) for FCSO vs. HCHC using restricted samples. The results were as follows: males only (i.e., females excluded) 0.69 (0.54, 0.87); individuals who are less involved with the criminal justice system (i.e., number of prior arrests >75th percentile excluded) 0.75 (0.57, 0.99); participants first arraigned as an adult (i.e., juveniles excluded) 0.64 (0.46, 0.90); participants on pre-trial status at index jail stay (i.e., sentenced individuals excluded) 0.64 (0.49, 0.84); FCSO participants with documented receipt of buprenorphine only (i.e., naltrexone or unknown discharge MOUD excluded) 0.72 (0.57, 0.93).

3.3. Mortality

During the first year after release, approximately 3% of participants from each site died, 6 participants from FCSO (median time-to-death 287.5 days, IQR [201,311], and 8 from HCHC (median time-to-death 141.5, IQR [14,310]). Of the FCSO deaths, 2 were due to overdose (both between 9 and 12 months after release), 2 were unknown causes, and 2 resulted from injury or disease. Of the HCHC deaths, 5 were attributed to overdose (3 occurred within the first month after release and 2 between 9 and 12 months after release), 2 had unknown causes, and 1 other causes. After one year post-exit from jail, an additional 6 deaths occurred among HCHC participants (3 from overdose) and no additional deaths among FCSO participants. The larger sample size and longer record review time for HCHC may explain observing at least some of these additional deaths. The mean±SD age at death was 42.2 ± 13.2 for FCSO and 40.9 ± 11.6 for HCHC.

4. Discussion

4.1. Key findings

This natural experiment across two similar rural jails in Massachusetts found that, among incarcerated adults with opioid use disorder, offering buprenorphine in jail substantially reduced the risk of recidivism. Results from the unadjusted Cox proportional hazards model found a 29% reduction in risk of recidivism, which reduced further to 32% after adjusting for baseline history of interactions with the criminal justice system and index jail status.

Post-hoc analyses found that recidivism related to property was reduced, in keeping with the logical supposition that MOUD effectively managed the opioid use disorder, and thus reduced associated drug related property crime. Analyses also found no differences in violations of parole or probation, suggesting that differing rates of rearrest for technical violations and variation in community correctional practices did not explain the findings.

The few prior studies detected limited impact of MOUD on recidivism (Perry et al., 2015). Gordon and colleagues (2017) examined outcomes of urban incarcerated individuals randomized to treatment with buprenorphine versus a counseling-only comparison group, reporting no differences in criminal activity one year after jail exit.
Magura and colleagues (2009) randomized urban incarcerated individuals to buprenorphine or methadone, and reported no differences in arrests, crime, or incarceration three months after jail exit. These studies both focused on populations returning to large metropolitan areas. The current findings suggest that MOUD may have more substantial impact among persons leaving smaller jails who return to rural communities, although more research is needed.

### 4.2. Limitations and strengths

Study findings must be considered with its limitations. Findings derive from an observational natural experiment in mostly rural settings in Massachusetts in which one jail provided access to buprenorphine and the other did not. However, participants were not randomized, raising the possibility of selection bias, omitted variable bias, or confounding as explanations for these findings. We did not account for potential variation by site in OUD screening and assessment practices, opioid overdose education practices, linkage to MOUD after jail exit, community re-entry services, availability of MOUD or other services in the community, or other legal and health system practices that may have impacted outcomes. On the other hand, adjustment for baseline characteristics strengthened the findings, suggesting that the magnitude of effect might be conservative, and findings did not change in sensitivity analyses using models with restricted samples. Furthermore, findings from the predominantly white male population in these small rural county jails may not generalize to all incarcerated persons with OUD. Both jails are Massachusetts Justice Community Opioid Innovation Network sites, which offers future opportunities to replicate findings among a larger and more diverse sample (Evans et al., 2021). Also, recidivism outcomes are measured with administrative records. Administrative data enable measurement of outcomes on all participants, a key reason why these data are useful for assessing addiction treatment outcomes (Evans et al., 2010, 2019; Krebs et al., 2017), but they provide information only on those events that resulted in an incarceration, probation violation, or arraignment in Massachusetts. Finally, receipt of psychosocial treatment in these jails was not measured, so its potential effects could not be

**Table 3**

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Recidivism (any)</th>
<th>Incarceration</th>
<th>Probation violation</th>
<th>Arraignment (any)</th>
<th>Arraigned: Drug</th>
<th>Arraigned: Property</th>
<th>Arraigned: Violent</th>
<th>Arraigned: Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>County: Franklin (ref = Hampshire)</td>
<td>0.51 (0.35, 0.76)</td>
<td>0.37 (0.24, 0.58)</td>
<td>0.91 (0.55, 1.52)</td>
<td>0.67 (0.45, 0.99)</td>
<td>0.76 (0.45, 1.28)</td>
<td>0.39 (0.22, 0.69)</td>
<td>0.70 (0.38, 1.28)</td>
<td>0.79 (0.43, 1.44)</td>
</tr>
<tr>
<td># of prior incarcerations</td>
<td>1.06 (1.02, 1.10)</td>
<td>1.03 (0.99, 1.07)</td>
<td>0.99 (0.95, 1.04)</td>
<td>1.06 (1.02, 1.10)</td>
<td>1.05 (1.00, 1.09)</td>
<td>1.05 (1.01, 1.09)</td>
<td>1.04 (0.99, 1.09)</td>
<td>1.02 (0.97, 1.07)</td>
</tr>
<tr>
<td>Jail status: pre-trial (index, ref)</td>
<td>2.05 (1.35, 3.12)</td>
<td>2.24 (1.41, 3.56)</td>
<td>2.27 (1.23, 4.21)</td>
<td>1.26 (0.83, 1.90)</td>
<td>1.52 (0.87, 2.67)</td>
<td>0.96 (0.58, 1.62)</td>
<td>1.27 (0.68, 2.38)</td>
<td>1.06 (0.57, 1.97)</td>
</tr>
</tbody>
</table>

**Fig. 1.** Time from jail exit to 1st recidivism event survival curves. Cox proportional hazards model unadjusted hazard ratio (95% CI) 0.71 (0.56, 0.89), *p* = 0.003. Adjusted for number of prior incarcerations, index jail status is pre-trial vs. sentence HR 0.68 (0.53, 0.86), *p* = 0.001.
examined.

4.3. Conclusion and policy implications

This natural experiment found substantial reductions in post-release outcomes among a large sample of individuals with OUD who received buprenorphine in jail compared to those who did not receive MOUD.

After decades in which access to agonist treatments for OUD was limited in corrections, lawsuits and legislation have created momentum for jails and prisons to provide these lifesaving therapies. In Massachusetts, a legislative mandate caused seven jails, including the two examined in the current study, to expand agonist treatment options and related services in 2019. These jails provide all three FDA-approved types of MOUD during incarceration, and programming to connect individuals to MOUD in the community at jail exit. In addition to the demonstrated reduction in overdose mortality, the current study provides legislators and correctional officials with compelling evidence that agonist MOUD in jail will reduce recidivism. Since recidivism and reincarceration are costly, and the implementation costs associated with agonist treatment in jails are substantial, future work should examine the state and societal costs associated with MOUD in jail.

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None to declare.

CRediT authorship contribution statement

PF and EE conceptualized the study, obtained funding, developed the protocols, and supervised data collection and data analysis. DW conducted the analysis. EE drafted the manuscript. All provided comments and finalized and approved the submitted version.

Declaration of Competing Interest

No conflict declared.

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