# Statistical Approaches to Automated Groove Engraved Area Identification in 3D Bullet Land Scans

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#### Abstract

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#### TO DO:

- Abstract
- Time to calculate for BCP

## 1 Background

Forensic pattern analysis aims to address the same-source problem: whether two impressions were generated by the same object. One of the most significant aspects of the same-source problem in firearms analysis is the determination of whether two bullets were fired through the same gun barrel. The evidence used in visual pattern comparison of bullets are striation marks, which are engraved on the bullet by micro imperfections in the barrel.

For barrels with traditional (i.e., not polygonal) rifling, striation marks found on land engraved areas (LEAs) are the primary evidence used. LEAs bear marks engraved by alternating sections of the barrel, and are considered to be areas which will bear unique patterns of striation marks which are reproduced on any bullet fired through the same barrel (1). Figure 1 depicts lands inside a barrel, as well as land engraved areas on a fired bullet.

The recent application of high resolution 3D scanning technology to bullet LEAs coupled with concerns about the objectivity of visual pattern comparison have motivated the development of several image-analysis algorithms which aim to complete automated, quantitative analyses of bullet evidence (see 2; 3; 4; 5; 6; 7; 8). The data used in such algorithms are high resolution 3D scans of LEAs, such as that pictured in Figure 2.

Bullets can be algorithmically compared to one another by completing individual LEA-to-LEA image comparisons of each LEA scan from one bullet to each LEA scan from a second bullet. Each LEA-to-LEA comparison compares the patterns of striation marks engraved on the LEA, which can be extracted from a 3D LEA scan as a 2D signature which details a pattern of peaks and valleys representing striation marks.

Signatures are extracted by first obtaining a horizontal slice of the 3D scan, called a profile, followed by removing the overall bullet curvature which dominates the data structure of each profile. Figure 3 depicts the process of translating a 3D LEA scan to a 2D

#### LEA signature.

Currently accepted best practice for capturing 3D LEA scans involves capturing portions of the neighboring groove engraved areas (GEAs), which introduce a secondary data structure on the edges of the LEA scan, which is not attributed to the pattern of interest: striation marks left on the LEA.

In order to accurately represent the striation pattern as a 2D LEA signature, the extraneous data structure introduced by GEA data must be identified and removed. The identification of GEA data, while quite straightforward for the human visual system, is a difficult process to automate using computer vision techniques.

We describe here two methods for automated identification and removal of GEA data. First, we use an adapted version of a robust statistical modeling technique to remove bullet curvature. Then, we describe two methods to separate GEA data from LEA data using statistical techniques. We then compare performance of the two approaches on three separate test sets of bullets.

## 2 Data Source

We use high resolution 3D scans of bullet LEAs. Scans were captured at Iowa State University's High Resolution Microscopy Facility on a Sensofar confocal light microscope at 20x magnification resulting in a resolution of 0.645 microns per pixel. Scans are stored as x3p files, conforming to the ISO5436-2 standard (9). x3p is the industry standard format for capture and storage of 3D microscopic topography of bullets. Objects are stored digitally as a 2-dimensional matrix with (x, y) locations corresponding to locations on the physical object; a relative height value z is measured and recorded for each (x, y) location.

All six LEAs from each bullet were captured for bullets from three separate test sets.

Hamby set 44 consists of 35 bullets fired from 10 consecutively rifled Ruger P85 barrels. There are two known bullets for each of the ten barrels, as well as 15 additional questioned bullets. Each fired bullet in Hamby Set 44 has 6 LEAs; every LEA was scanned for each of the 35 bullets, producing data for 210 individual land engraved areas. Two lands – Barrel 9, Bullet 2, Land 3 and Unknowns, Bullet L, Land 5 – were removed from consideration due to "tank rash". Tank rash results from a bullet striking the bottom of a water recovery

tank after exiting the barrel, thereby creating marks on the land that are not due to the contact with the barrel.

The Phoenix PD set consists of 33 bullets fired from 8 barrels more information on the type of barrels?. There are three known bullets for each of the eight barrels, as well as 9 additional questioned bullets. There are 6 LEAs for each of the fired bullets, producing a total of 198 individual land engraved areas.

The Houston-test set consists of 45 bullets fired from at least 10 barrels more information on the type of barrels?. There are three known bullets for each of the ten barrels, as well as 24 questioned bullets that are fired from a combination of the 10 known barrels and additional, out-of-set barrels. There are 6 LEAs for each of the fired bullets, producing a total of 414 individual land engraved areas.

A crosscut was extracted from each scan by identifying an optimal crosscut using  $x3p\_crosscut\_optimize$  in the bulletxtrctr package in R (10). We calculated an averaged profile by averaging across ten consecutive crosscuts which fall directly to either side of the optimal crosscut identified. The following GEA data identification methods are applied to these averaged profiles, for a total of 820 profiles across the three test sets. In the following, we utilize relative horizontal locations on the profile,  $x_i$ , and their relationship with relative height values on the profile,  $z_i$ , for data points i = 1, ..., n on each individual profile.

## 3 Methodology

#### 3.1 Global Structure Removal

The first stage of our GEA removal process focuses on removing the bullet curvature from each profile. GEAs on the edge of each LEA represent a change in the primary data structure typically characterized by a sharp increase in measured height values, as demonstrated in Figure 3(Middle). Removal of the primary curvature should leave the secondary structure intact – namely, the sharp increase in height values. The increase in values can then be used to separate LEA data from GEA data.

Bullets are subjected to significant pressure in the process of being fired through a gun

barrel. Thus, we cannot assume completely circular curvature on bullet LEAs. Further, we cannot model bullet curvature as a directly circular structure. We use more flexible non-parametric locally weighted regression (LOESS), which can model large-scale structure of the height values  $z_i$ . LOESS is flexible enough to model structural defects within the bullet curvature. However, the flexibility also leaves LOESS susceptible to modeling the secondary structure of the GEA.

Traditional LOESS predicts height values  $\hat{z}_i$  as a function of location  $x_i$  by estimating values  $\beta_0, \beta_1$  which minimize:

$$\arg\min_{\beta} \sum_{k=1}^{n} w_k(x_i) (z_k - (\beta_0 + \beta_1 x_k))^2,$$

where  $w_k(x_i)$  is a weight assigned to each data point  $x_k$  based on its proximity to  $x_i$ . Weights  $w_k$  decrease as distance to  $x_i$  increases, so that data points closest to  $x_i$  influence the prediction  $\hat{z}_i$  most. Weights  $w_k(x_i)$  are defined using a prespecified decreasing function, traditionally a tricube weight.

An approach which mitigates the impact of GEA data is robust LOESS (see (11)), a process which iteratively updates weights  $w_k(x_i)$  by multiplying by a robustness weight,  $\delta_k$  based on the magnitude of each residual  $e_k = z_k - \hat{z}_k$ . Larger values of  $e_k$  result in a lower weight for that data point,  $z_k$ . The process is as follows:

- 1. Fit a LOESS model to a LEA profile, predicting height  $z_i$  using relative location  $x_i$ . Assign robustness weights  $\delta_k$  of 1 to each data point  $(x_k, y_k)$ .
- 2. Obtain predicted height values  $\hat{z}_k$ , and corresponding residual values  $e_k = z_k \hat{z}_k$ .
- 3. Calculate updated robustness weights using residual values  $e_k$ :

$$\delta_k \times w_k(x_i) = \left(1 - \left(\frac{e_k}{6*MAD}\right)^2\right)^2 \times w_k(x_i) \quad \text{if} \quad \left|\frac{e_k}{6*MAD}\right| < 1,$$

where MAD is the median absolute deviation of residuals  $e_k$ . This is known as the bisquare function.

4. Repeat steps 1-3 with updated weights at each iteration for m iterations, with 20 iterations as the default.

5. After m iterations of updating the weight vector  $\delta_k w_k(x_i)$ , fit a LOESS model and obtain residual values  $e_i$  for each data point  $(x_i, z_i)$ .

Re-weighting data as in Step 3 reduces influence of data points with large absolute residual values. When GEA data is present on a profile, largest residual values will occur in areas where GEA data begins as it presents a competing structure with overall LEA curvature.

Figure 5 depicts the impact this iterative re-weighting process has on curvature removal for an example LEA from Hamby set 44. Predictions  $\hat{z}_i$  are updated and more closely follow the primary structure of bullet curvature.

However, this method fails to mitigate GEA data impact when GEA structures are more pronounced (see Figure 6). Thus, we make a small adaptation to the procedure to function more effectively with these data structures:

- 1. Fit a LOESS model to a LEA profile, predicting height  $z_i$  using relative location  $x_i$ . Assign robustness weights  $\delta_k$  of 1 to each data point  $(x_k, y_k)$ .
- 2. Obtain predicted height values  $\hat{z}_k$ , and corresponding residual values  $e_k = z_k \hat{z}_k$ .
- 3. Calculate updated robustness weights using residual values  $e_k$ :

$$\delta_k \times w_k(x_i) = \left(1 - \left(\frac{e_k}{6*MAD}\right)^2\right)^2 \times w_k(x_i) \quad \text{if} \quad \left|\frac{e_k}{6*MAD}\right| < 1,$$

where MAD is the median absolute deviation of residuals  $e_k$ .  $\delta_k$  is known as the bisquare function.

- 4. Assign updated weights as in Step 3 if  $e_k > 0$ . Else, leave weights as  $w_k(x_i)$ .
- 5. Repeat steps 1-3 with updated weights at each iteration for m iterations, with 20 iterations as the default.
- 6. After m iterations of updating the weight vector  $w_k(x_i)$ , fit a LOESS model and obtain residual values  $e_i$  for each data point  $(x_i, z_i)$ .

This adapted robust LOESS, while a small procedural change, more adeptly fits bullet curvature. Figure 7 demonstrates the impact of the adaptation on predicted height values  $z_i$ . This impact differs for each of the bullet test sets. Figure 8 depicts the differing levels of impact our adjustment has, dependent on bullet test set. Once I have more barrel information, I could go into a little more detail here about differing barrel types/ammo types or something, but no more than a sentence to bring the point home.

Once the adapted robust LOESS procedure has been applied to a LEA profile, the resulting residuals  $e_i$  follow a reliable pattern: small residuals closer to zero in locations associated with LEA data, and sharply increasing, larger residuals in locations associated with GEA data. The resulting residual structure lends itself to statistical techniques to separate the two structures more effectively.

The subsequent GEA identification methods are based on residuals  $e_i$  calculated from adapted robust LOESS fit to the global structure of the profile. One method, using supervised two-class classification techniques, aims to classify each data point as being part of the LEA or GEA. The second approach is an unsupervised method based on change-point analysis that seeks to identify data points where the distribution of the residuals changes. Both methods result in "shoulder location predictions", predictions for the locations  $(x_L, x_R)$  at which LEA data ends and GEA data begins on the left and right side of the profile, respectively.

#### 3.2 Two-Class Classification

Shoulder location can be predicted by first classifying data points as one of two classes ("LEA" or "GEA"), and subsequently gathering the range of values classified as "LEA" points.

Classification into "LEA" or "GEA" is first approached by a process of feature engineering based on adapted robust LOESS residuals. While the residuals  $e_i$  should demonstrate differing patterns of magnitude, residuals alone are not enough to classify data with high accuracy.

Figure 9 demonstrates that each test set has distinct patterns of striation depth. Thus, standardization of features is imperative for transferability of fitted model parameters. The

LEA scan context requires some non-traditional standardization practices.

For example, consider the distribution of residual values  $e_i$  resulting from adapted robust LOESS. There is reason to believe that the distribution will be quite skewed, which means a standard deviation will not be a good proxy for the spread of the distribution. Thus, rather than the standard deviation, we consider instead the standard deviation of residual values from the middle 50% of  $x_i$  values present in each profile. This alternative acts as a proxy for the depth of striae on each LEA, with higher standard deviations found for lands with deeper striae. Standardizing residual values by this proxy puts all residuals on a comparable scale.

For variables which instead deal with differences in the x direction, such as depth from the center of a scan, values will be mapped to a (0,1) range, with the maximum x location on the scan acting as the divisor.

The full list of standardized features are as follows:

rlo\_resid\_std: Robust LOESS residual value  $e_i$ , standardized by dividing by standard deviation of residual values from middle 50% of  $\mathbf{x_i}$  values.

 $(rlo\_resid\_std)^2$ : Squared term of  $rlo\_resid\_std$ .

side: Whether data point is to left or right of median  $x_i$  value.

depth\_std: Distance of data point from median  $x_i$  value, standardized by dividing by maximum  $x_i$  value (a proxy for the range of x values).

side:depth\_std: Interaction between side and depth\_std variables.

xint1\_std: Predicted location  $x_i$  at which adapted robust LOESS crosses y axis on left side of profile, standardized by dividing by maximum  $x_i$  value (a proxy for the range of x values).

xint2\_std: Predicted location  $x_i$  at which adapted robust LOESS crosses y axis on right side of profile, standardized by dividing by maximum  $x_i$  value (a proxy for the range of x values).

range\_50\_std: Range of residual values  $e_i$  within a 50-point window  $(x_{i-25}, x_{i+25})$  around data point  $x_i$ , standardized by dividing by standard deviation of residual values from middle 50% of  $x_i$  values.

numNA\_50: Number of missing values within a 50-point window  $(x_{i-25}, x_{i+25})$  around data point  $x_i$ .

ind\_2mad: Indicator of whether rlo\_resid is greater than  $2*MAD(rlo_resid)$ , where MAD is the median absolute deviation of residual values  $e_i$  for an entire profile.

numpos\_50: Number of positive residual values within a 50-point window  $(x_{i-25}, x_{i+25})$  around data point  $x_i$ .

ind\_edges: Indicator of whether data point is to the left of xint1 or to the right of xint2. Values between xint1 and xint2 receive a value of 0, while values on the outside of the two values receive a value of 1.

Examples of the distributions of some of these features can be seen in Figure 10.

A logistic LASSO model, a form of penalized regression, was fit using the developed features. LASSO parameter values for p covariates were calculated by identifying:

$$\hat{\beta}_{\lambda} = \beta \in \mathbb{R}^{p} \left\{ (Y - X\beta)'(Y - X\beta) + \lambda \sum_{j=1}^{p} |\beta_{j}| \right\}$$

which adds a penalty to the traditional ordinary least squares minimization problem, and uses a tuning parameter  $\lambda$  (12).

A cross-validated LASSO model was fit using the cv.glmnet function in the glmnet package in R (13). Parameter values from the model with  $\lambda_{1se}$  were used.  $\lambda_{1se}$ , a standard when using LASSO, is the tuning parameter which results in the simplest model that still has cross-validation error within one standard deviation of the best model.

The resulting model, "LASSO Full", uses each of the data features listed above along with pairwise interactions for each of them. These parameter values were trained on the Hamby set 44 data. Parameter values were used to calculate predicted values of GEA membership between 0 and 1; the closer to 1, the higher probability of membership in the "GEA" class.

Two-class classification techniques traditionally employ a cutoff for predicted probabilities and assign predicted class membership using that cutoff; i.e., values above a certain cutoff are classified as part of the "GEA" class, and values below the cutoff are classified as part of the "LEA" class. An equal error rate is typically used for this purpose. However, since LEA scans consist of a majority of LEA data by nature, significant class imbalance in our response variable ("GEA" vs. "LEA") dictates a change in procedure for assigning a cutoff for class membership. Equal error rate is defined as the cutoff at which sensitivity (true positive rate) and specificity (true negative rate) are equal. Employing this rate to predict class membership would result in a higher number of false positives, here predicting data which is part of the "LEA" class as "GEA" data. We employ an equal number of errors rate rather than an equal error rate to ameliorate this imbalance.

The "LASSO Full" method for shoulder location identification is as follows:

- 1. Use adapted robust LOESS procedure to remove bullet curvature and obtain residual values  $e_i$ .
- 2. Extract data features based on  $x_i$  locations and residual height values  $e_i$  from Step 1.
- 3. Use fit parameter values from pre-fit LASSO model to calculate probabilities of membership in GEA class.
- 4. Apply cutoff of .34: classify higher probabilities as GEA data points, and lower probabilities as LEA data points.
- 5. Identify the minimum  $x_i$  value which is predicted as a member of the LEA class,  $x_L$ . Identify the maximum  $x_i$  value which is predicted as a member of the LEA class,  $x_R$ .  $(x_L, x_R)$  are the shoulder location predictions.

This method for shoulder location identification can be found in the R package bulletxtrctr as the function get\_grooves\_lassofull.

### 3.3 Bayesian Changepoint Analysis

Our second approach is also based on the idea that residuals  $e_i$  resulting from adapted robust LOESS predictions will follow a consistent pattern: decreasing values in the left GEA, values with no discernable slope change in the LEA, and increasing values in the right GEA. This can be thought of as a line with negative slope for the left GEA, line with zero slope for the LEA, and a line with positive slope for the right GEA. Our model will be therefore be defined in a piecewise fashion. The points of global structural change are what we will call changepoints. Changepoint locations can be treated as parameters and estimated in the same manner as any other parameter in a statistical model.

This approach was proposed more generally as Bayesian changepoint detection in Stephens (14). In the context of LEA residuals, there are additional complex patterns (i.e., striation marks), but the overall linear structure remains on a larger scale. We will consider the smaller scale striation patterns as dependence in the data after accounting for the large scale structures we are estimating.

We first perform a few additional data processing steps to streamline computation. First, we scale the residuals  $e_i$  from the robust LOESS procedure by dividing by the standard deviation of residuals  $e_i$  for the entire profile. Secondly, we impute missing residual values  $e_i$  to maintain equidistance between  $x_i$  locations.

We next develop a model which we will use to identify changepoints, and describe estimation procedures. Additional details on data processing can be found in the appendix.

#### 3.3.1 Bayesian Model Formulation

For model formulation, we will define random variables associated with residual values  $e_i$ . Let  $\{Y(x_i): i=1,2,...,n\}$  denote the set of random variables representing the residuals from the robust LOESS procedure at the values  $x_i$ . For simplicity, also assume that  $x_1 < x_2 < ... < x_n$ . Also, let  $c_l$  be the value of the left changepoint and  $c_r$  be the value of the right changepoint. Here, the left changepoint is where the left GEA meets the LEA, and the right changepoint is where the right GEA meets the LEA. Also, denote the median centered x values as  $x'_i = x_i - \tilde{x}$  where  $\tilde{x}$  is the median x value. As mentioned in the previous paragraph, the complex small scale patterns, such as the striae, will be modeled

through a covariance structure on the data that will be allowed to differ between each GEA and between the GEAs and LEA. We will construct the covariance matrices from the exponential covariance function  $K(x, x'; \sigma, \ell) = \sigma^2 e^{-\frac{|x-x'|}{\ell}} = cov(Y(x), Y(x'))$ . The differences in covariance matrices for the GEAs and LEA will be reflected in the parameters  $\sigma$  and  $\ell$ . The data model that we consider is then,

$$(Y(x_1), Y(x_2), ..., Y(x_{k_1}))^{\top} \sim N(\beta_{01} \mathbb{1} + \beta_{11} x_{1:k_1}, \Sigma_1(\sigma_1, \ell_1))$$
 (1)

$$(Y(x_{k_1+1}), Y(x_{k_1+2}), ..., Y(x_{k_2}))^{\top} \sim N(0, \Sigma_2(\sigma_2, \ell_2))$$
 (2)

$$(Y(x_{k_2+1}), Y(x_{k_2+2}), ..., Y(x_n))^{\top} \sim N(\beta_{02} \mathbb{1} + \beta_{12} x_{k_2+1:n}, \Sigma_3(\sigma_3, \ell_3)),$$
 (3)

where  $x_{k_1} < c_l \le x_{k_1+1}$  and  $x_{k_2} < c_r \le x_{k_2+1}$  Here,  $x_{1:k}$  denotes the column vector  $(x_1, x_2, ..., x_k)^{\top}$ , and  $\mathbbm{1}$  denotes the vector of ones. Independence is assumed between each of these three distributions for simplicity. The parameters that need to be estimated include the four mean parameters in the GEAs, the six covariance parameters (two for each of the three areas), and the two changepoint parameters,  $c_l$  and  $c_r$ .

The above model encapsulates the essence of the approach. However, there are a few difficulties. The first difficulty is that there are not always two GEAs in a particular land. There may be one GEA, or the land may only consist of the LEA. Thus, the above model is actually conditional on there being two GEAs in the data. We also define models for when there is one GEA on the left, one GEA on the right, or no GEAs. The models are defined in an essentially identical way. Conditional on there being only one GEA, the left GEA model is defined as,

$$(Y(x_1), Y(x_2), ..., Y(x_k))^{\top} \sim N(\beta_0 \mathbb{1} + \beta_1 x_{1:k}, \Sigma_1(\sigma_1, \ell_1))$$
 (4)

$$(Y(x_{k+1}), Y(x_{k+2}), ..., Y(x_n))^{\top} \sim N(0, \Sigma_2(\sigma_2, \ell_2)),$$
 (5)

and the right GEA model is defined as,

$$(Y(x_1), Y(x_2), ..., Y(x_k))^{\top} \sim N(0, \Sigma_1(\sigma_1, \ell_1))$$
 (6)

$$(Y(x_{k+1}), Y(x_{k+2}), ..., Y(x_n))^{\top} \sim N(\beta_0 \mathbb{1} + \beta_1 x_{k+1:n} \Sigma_2(\sigma_2, \ell_2)).$$
 (7)

Finally, conditional on there being no GEAs in the data, the model is simply

$$(Y(x_1), Y(x_2), ..., Y(x_n))^{\top} \sim N(0, \Sigma(\sigma, \ell)).$$
 (8)

We see that estimating the changepoint locations also involves selecting the most appropriate model from these four. In order to avoid confusion, we have slightly abused notation and, for example,  $\Sigma_1(\sigma_1, \ell_1)$  as it is estimated in the two changepoint model is *not* the same as  $\Sigma_1(\sigma_1, \ell_1)$  from either of the one changepoint models, and  $\Sigma_1(\sigma_1, \ell_1)$  is also *not* the same between the two one changepoint models. As another example,  $\beta_0$  is *not* the same between each of the one changepoint models. So, to be clear, duplication of notation in different models is not meant to imply that those parameters are shared between models.

Ultimately, these above four models are each individually fitted, and each model above is given a prior. From there, we do model selection in the formal Bayesian way, selecting number and location of changepoints by maximizing the estimated posterior distribution.

In order to complete a Bayesian model specification, we need priors on each of the parameters in each model as well as each model itself. We will assume independence between each parameter a priori. For each length scale  $\ell$ , we will assume  $\ell \sim \text{Gamma}(3,5)$ . For each standard deviation, we will assume  $\sigma \sim \text{Half-Normal}^+(0,1)$ , where  $\text{Half-Normal}^+(\cdot,\cdot)$ is notation for the normal distribution restricted to the positive real numbers. For intercept parameters,  $\beta_{01}, \beta_{02}, \beta_0 \sim N(0, 10)$ . For the slope parameters, the preceding trend deviates slightly. For any slope that corresponds to the left GEA,  $\beta_1$  or  $\beta_{01}$ , we will assume that the slope can not be positive. That is,  $\beta_1, \beta_{01} \sim \text{Half-Normal}^-(0, 10)$ , where Half-Normal $^-(\cdot, \cdot)$ is notation for the normal distribution restricted to the negative real numbers. Contrastingly, for any slope that corresponds to the right GEA,  $\beta_1$  or  $\beta_{02}$ , we will assume that the slope can not be negative. That is,  $\beta_1, \beta_{01} \sim \text{Half-Normal}^+(0, 10)$ . For the changepoint locations, we assume a uniform prior  $\pi(c_l, c_r) \propto I(a < c_l < c_r - \gamma < b - \gamma)$ . Here, a and b are some values close to the edges of the data. How close those values are to the edges is a parameter that is set manually. Further, we include another hyperparameter,  $\gamma$ , which can be set so that the changepoints are not allowed to be too close to each other. This is also a parameter that is set manually. Lastly, we assume a uniform prior over all four models.

#### 3.3.2 Bayesian Model Estimation

As was noted in Stephens (14), for any model including a changepoint, the likelihood is not a smooth function of the changepoint location. This is because, holding all other parameters fixed, shifting the changepoint value will result in zero change to the likelihood until it crosses the nearest point to the right or left, at which point the likelihood makes a jump. This makes maximum likelihood estimation in the standard way infeasible, but Bayesian estimation can be done in a fairly straightforward way via Markov chain Monte Carlo (MCMC). The basic idea is that, for each model, we can construct a two step Gibbs sampler. In step 1 we sample from the posterior distribution of the mean and covariance parameters given the changepoint locations, and in step 2 we sample from the changepoint locations given the mean and covariance parameters. Because of the non-conjugacy in our model, we perform both sampling steps using a random walk Metropolis-Hastings (RWMH) step with Gaussian proposals. For details on Gibbs sampling and the Metropolis-Hastings algorithm see Gelman et al. (15). It is also worth mentioning that the zero changepoint model does not require Gibbs sampling at all, and we perform estimation there using a RWMH algorithm.

We now provide the two basic steps of the Gibbs sampler for the two changepoint case. The algorithms to sample from the other three models are omitted, and are nearly identical except for the smaller number of parameters that need to be sampled. Denote collection of mean and covariance parameters for the left GEA as  $\theta_1$ , the LEA as  $\theta_2$ , and the right GEA as  $\theta_3$ . Then, at iteration t after warmup

- 1. given changepoint locations  $(c_l^{(t-1)}, c_r^{(t-1)})$ , sample  $(\theta_1^{(t)}, \theta_2^{(t)}, \theta_3^{(t)})$  using independent RWMH steps for each  $\theta_i$
- 2. given  $(\theta_1^{(t)}, \theta_2^{(t)}, \theta_3^{(t)})$ , sample  $(c_l^{(t)}, c_r^{(t)})$  using a single RWMH step.

After running the MCMC for each model, parameter estimates and the most likely model are jointly chosen according to the largest joint posterior value. That is, we arrive at estimates  $(\hat{\theta}, \hat{M}) = \underset{(\theta, M)}{\operatorname{argmaxlog}}(p(\theta, M|Y))$ , where M is the random variable associated with the choice of model,  $\theta$  is the associated parameter vector for the appropriate model, and Y is all of the available data. Additional MCMC details can be found in the appendix.

### 4 Results

To assess the degree of improvement in automated shoulder location identification, we want to quantify the impact each prediction method has on an automated bullet matching algorithm's accuracy. Four different shoulder location identification methods will be inserted into the automated bullet matching algorithm process:

- (1) rollapply, the method proposed by (8),
- (2) LASSO full,
- (3) Bayesian changepoint, and
- (4) manual identifications, the "gold standard" for identification.

Shoulder location predictions were found using each of these methods, and used to remove GEA data from each profile. This is followed by extracting a signature, and using extracted signatures for each land engraved area to calculate pairwise similarity scores for all LEA signatures within each test set. Pairwise similarity scores were calculated using the random forest algorithm in the bulletxtrctr package.

Random forest scores should be close to 1 for land-to-land comparisons between signatures that originate from the same land, and closer to 0 for land-to-land comparisons between signatures from different lands.

We will investigate these scores for each individual test set: Hamby set 44, Phoenix PD set, and Houston-test set. All pairwise comparisons within a test set were completed. We look at both visual representations of the random forest score distributions as well as investigate the random forest method's accuracy in determining whether two lands originate from the same source.

There should be visual separation between "same source" and "different source" distributions, as is present for Manual ID predictions for all three test sets, seen in Figure 11, Figure 12, and Figure 13. LASSO full and Bayesian changepoint both show improvement in separation for all three test sets; however, there is still room for improvement when compared to Manual ID distributions in the Phoenix PD and Houston-test sets.

Quantitatively, there is also significant improvement in AUC values for all three test sets, as seen in Table 1, Table 2, and Table 3. Of interest is the classification accuracy with a fixed false positive rate. Since false positives - in this case identifying two land engraved areas as same source when they are different source - are the worst possible mistake in the forensic context, we set a cutoff rate for random forest scores based on a controlled false positive rate of .01.

Accuracy is overall high regardless of shoulder location identification method; however, there was a significant reduction in number of false negatives for the LASSO full and Bayesian changepoint methods for both Hamby set 44 and Phoenix PD sets. For the Houston-test set, Bayesian changepoint improves upon rollapply, but lags behind the LASSO full method in AUC and reduction of false negatives.

## 5 Conclusions

Both proposed approaches show significant improvement both visually and quantitatively over the "rollapply" method proposed by (8). However, the LASSO full method shows greater improvement than Bayesian changepoint on the Houston-test set. While manual identification of shoulder locations is still the most accurate method, and considered the "gold standard", the reduction in time to get predictions when using LASSO methods is advantageous and allows for less human involvement in the overall automated matching process.

The initial appeal of unsupervised methods such as Bayesian changepoint was the lack of dependence on training data and hence, potential generalizability advantages. This advantage did not bear out in the test sets presented here, and it appears the LASSO method trained on Hamby set 44 generalized effectively to new data.

While improvement is apparent on all three test sets using the LASSO and Bayesian changepoint methods, there is clear room for additional precision. Future work on the LASSO method should include re-training the LASSO model on a wider variety of LEA types rather than just the Hamby set 44 to avoid over-fitting to a specific type of LEA.

## 6 Appendix

#### 6.1 MCMC Details

As a practical note, it turns out that the posterior distribution is almost always multimodal, and it can happen that the sampler gets stuck in a suboptimal mode for a large number of iterations. It is also the case that the suboptimal modes need not even be close to the groove locations. It has, however, been our experience that the optimal mode corresponds well to the actual groove locations, which are often somewhat close to the edges of the data. With this in mind, starting values and the RWMH proposal variances play a very important role in the success of the sampling algorithm. Fortunately, it seems to be the case that by setting the initial changepoint values close to the edges of the data and making the proposal variance small (around 100 seems to work well) allows the sampler to wander inwards, and even with a modest number of iterations (say 5000), typically pass through the largest mode corresponding to the groove locations. This is not always the case, and it is possible that increasing the number of iterations produces better results.

In our implementation of this algorithm, the sampling functions were originally written with the intention of tuning the proposal variances to potentially accelerate convergence, and thus several warmup iterations are required for this purpose. This turns out to be a bad idea in this context for two reasons. The first reason is that the warmup iterations allow the sampler to wander past the global modes and get stuck in suboptimal modes far from the groove locations, from which the sampler may or may not find its way back to the optimal modes in just a few thousand iterations. Secondly, if the sampler does wander past the optimal modes, which are usually on the edges of the data, the tuned proposal variance can be quite large. The large proposal variance might not be a huge problem if it weren't for the fact that the width of the modes are almost always quite small. This means that it can take a very, very long time for the sampler to move from a suboptimal mode to the global mode. In order to mitigate this problem, we are currently setting the number of warmup iterations to be relatively small (somewhere in 100 to 500 seems to work well). In future, our implementation of the algorithm will not require any warmup iterations.

Initially, the Metropolis proposal variance for each  $\theta_i$  is diagonal with diagonal elements

all equal to 1/2. The proposal variance for  $(c_l, c_r)$  is initially set to be diagonal with elements equal to  $10^2$ . Note that because of the currently necessary warmup iterations, the variances after warmup for each  $\theta_i$  becomes  $\frac{2.4^2}{d}\hat{Var}(\theta_i^{(1:w)}) + \text{diag}(0.1)$ , where d is the dimension of  $\theta_i$  (which is not constant between GEAs and LEA), and  $\hat{Var}(\theta_i^{(1:w)})$  is the estimated variance covariance matrix from the w warmup iterations. Note that the addition of a diagonal matrix with entries 0.1 is to avoid the case when most or all warmup iterations have the same value. Similarly, the proposal variance for  $(c_l, c_r)$  after warmup becomes  $\frac{2.4^2}{2}\hat{Var}((c_l, c_r)^{(1:w)}) + \text{diag}(1)$ .

### 6.2 Data Preprocessing for MCMC

Before running the MCMC to do the changepoint detection, we first perform two data preprocessing steps. The first step is to scale the residuals from the robust loess procedure by the standard deviation calculated from the entire set of residuals. The reason for this is simply to make priors for standard deviation and slope parameters easier to specify. For example, ensuring that the residuals are scaled to have standard deviation one means that the standard deviation parameters in our model should also be close to one. This scaling also ensures that slopes values are not very large.

The second preprocessing step is a bit more involved. In order to enable the algorithm to run reasonably fast, we need to take advantage of the sparse precision matrix structure that is induced by the exponential covariance function. Indeed, this was the reason for choosing this covariance function in the first place. Unfortunately, it is challenging to do this unless the observations are evenly spaced in the domain. In our case, this would be true if there were no missing values. In order to remedy this problem, we impute the missing data, but only in the case that there exist non-missing observations outside of the missing values. In the case that the missing values exist on the edges of the data, we simply do not consider those domain values in the model.

We perform the imputation by treating the observations as coming from an unknown function, and infer the missing values from the known function values. In order to do this, we model the data with a Gaussian process and the squared exponential covariance function. That is, we suppose that

$$Y(x) \sim \mathcal{GP}(0, K(x, x'; \sigma^2, \ell)),$$

where now  $K(x, x'; \sigma^2, \ell) = \sigma^2 e^{-(x-x')^2/(2\ell^2)}$  is the squared exponential covariance function. We emphasize for clarity that this is a different covariance function than we use in the changepoint model. The main reason for this is that in imputing values, it seems desirable to allow dependencies beyond immediately neighboring points to influence predictions as the function that we are trying to predict generally has a smooth global structure. For all of our experiments, we set  $\sigma = 0.8$  and  $\ell = 15$ . These values were chosen from doing maximum likelihood estimation for a representative bullet.

When we impute the missing values, we compute the conditional mean of the missing values. To be clear, denote the distribution of the observed and missing data as

$$(Y, Y^*)^{\top} \sim N \left( \begin{bmatrix} 0 \\ 0 \end{bmatrix}, \begin{bmatrix} \Sigma_{yy} & \Sigma_{yy^*} \\ \Sigma_{y^*y} & \Sigma_{y^*y^*} \end{bmatrix} \right).$$

Here, Y is observed data and  $Y^*$  is the missing data, and the covariance matrix above is constructed from the squared exponential covariance function. We then use normal distribution theory to calculate the imputed values

$$E(Y^*|Y=y) = \sum_{y^*y} \sum_{yy}^{-1} y$$

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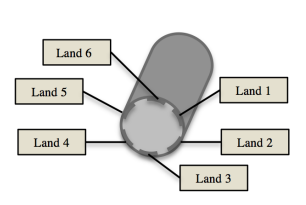




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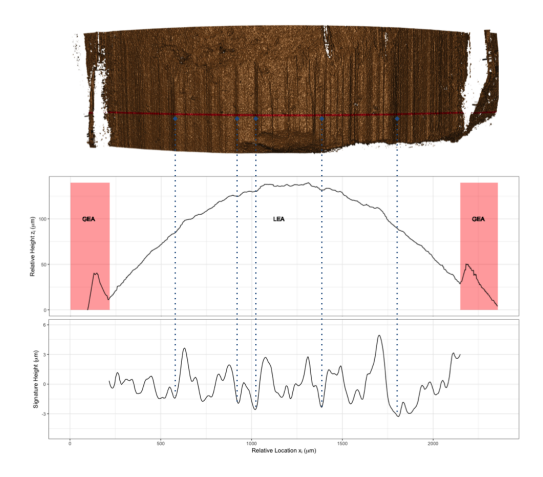


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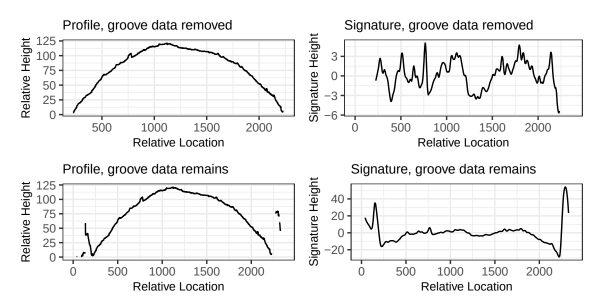


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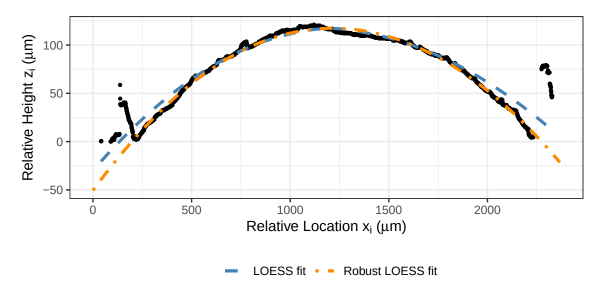


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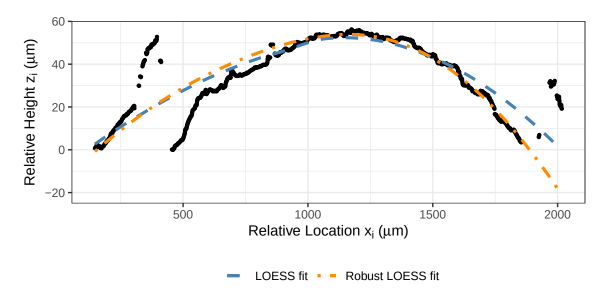


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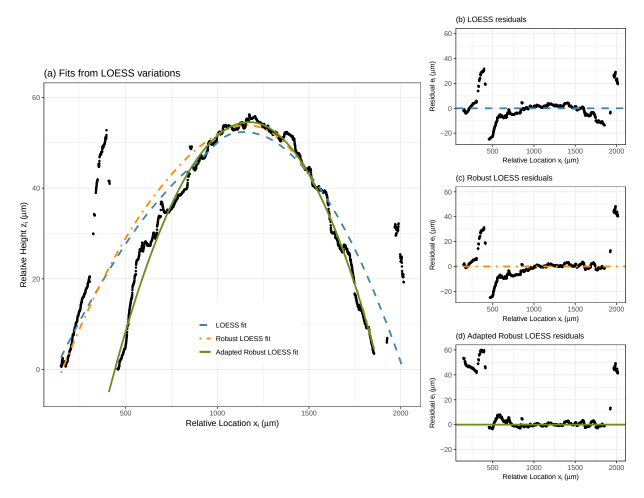
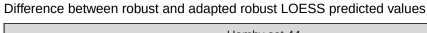


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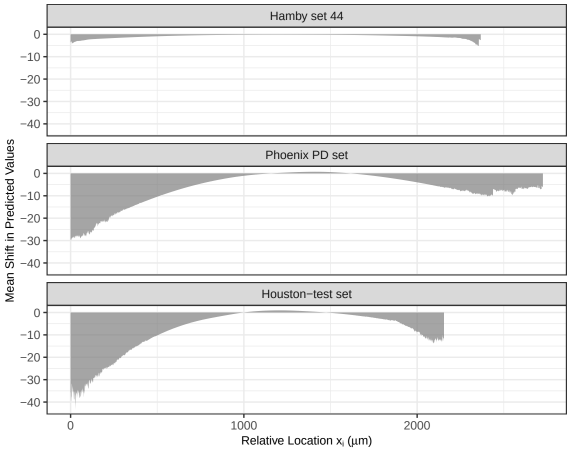


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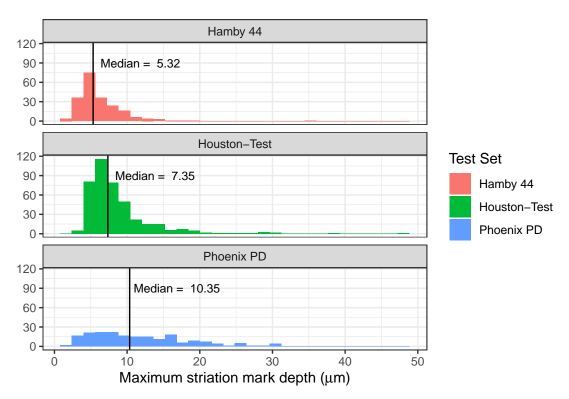


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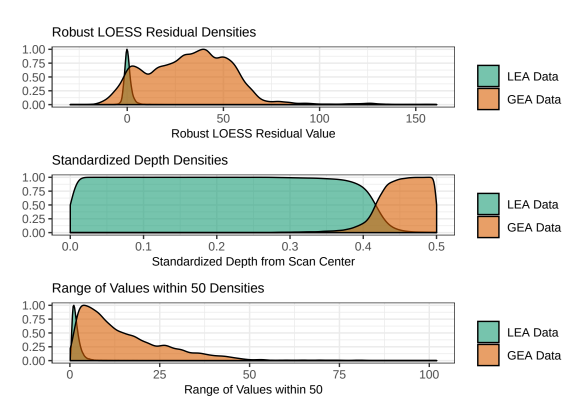


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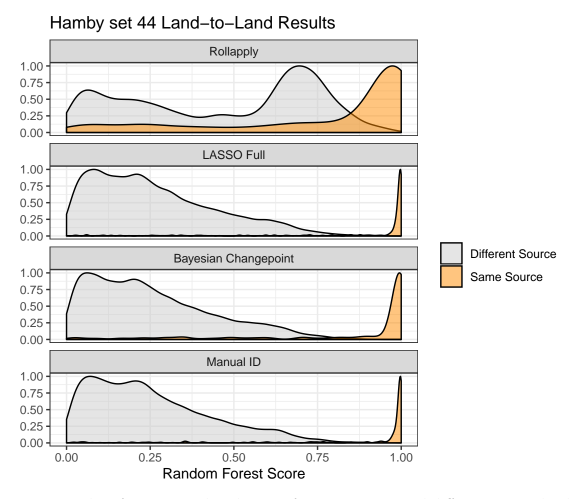


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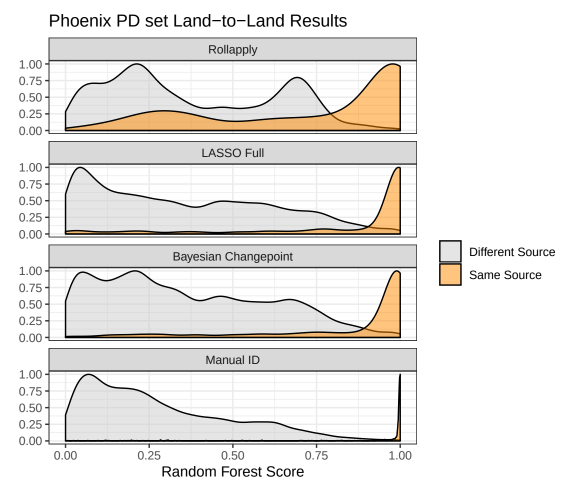


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## Houston-test set Land-to-Land Results Rollapply 1.00 0.75 0.50 0.25 0.00 LASSO Full 1.00 0.75 0.50 0.25 0.00 Different Source **Bayesian Changepoint** 1.00 Same Source 0.75 0.50 0.25 0.00 Manual ID 1.00 0.75

Figure 13: Random forest score distributions for same source and different source land-to-land comparisons for the Houston-test set. LASSO Full demonstrates improvement over Rollapply, but is still not as well separated as the Manual ID distributions. Bayesian changepoint demonstrates minor improvement, but does not improve as much as the LASSO method.

0.50

Random Forest Score

0.75

1.00

0.50 0.25 0.00

0.00

0.25

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# Controlled FPR = .01

Method	$\mathbf{AUC}$	Cutoff	FN	TP	TN	Accuracy	Time to Calculate
Rollapply	0.8	0.92	332	420	42126	0.98	1 min.
LASSO full	0.94	0.75	126	626	42106	0.99	6 min.
Bayesian Changepoint	0.93	0.74	152	600	42098	0.99	150 min.
Manual ID	0.94	0.74	124	628	42088	0.99	45 min.

Table 1: Land-to-land comparison results for Hamby set 44.

# Controlled FPR = .01

Method	$\mathbf{AUC}$	Cutoff	FN	TP	TN	Accuracy	Time to Calculate
Rollapply	0.818	0.9	364	378	38082	0.98	1 min.
LASSO full	0.893	0.953	238	504	38082	0.98	6 min.
Bayesian Changepoint	0.903	0.937	256	486	38080	0.98	140 min.
Manual ID	0.953	0.853	96	646	38084	0.99	45 min.

Table 2: Land-to-land comparison results for the Phoenix PD set.

## Controlled FPR = .01

Method	AUC	Cutoff	FN	TP	TN	Accuracy	Time to Calculate
Rollapply	0.761	0.91	1652	1016	167118	0.98	2 min.
LASSO full	0.858	0.823	1144	1524	167062	0.98	12 min.
Bayesian Changepoint	0.795	0.863	1552	1116	167096	0.98	280 min.
Manual ID	0.931	0.823	614	2054	167098	0.99	75 min.

Table 3: Land-to-land comparison results for the Houston-test set.