# "NEAREST-TREE" ESTIMATIONS 

# A discussion of their geometry 

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#### Abstract

The use of "nearest-neighbor" sampling has a long history. It involves measuring the distance from a random point in an area to the nearest object. That history involves never quite solving the problem, many examinations of special cases that never occur, adjustments that were ad-hoc, and a great deal of uninformative algebra. In forestry we have attempted to use the "nearest-tree" method for estimating numbers of trees on a landscape but the method is general, and can be used for any objects being sampled. I believe that the literature has never shown the logic and geometry in a form that is useful to both understand and solve the problem. This paper discusses the method from the geometric point of view, making no assumptions about tree distribution, and shows why extending the processes to the " n th closest tree" much reduces the bias and variability, as well as specifying what is needed to solve the problem in an unbiased way.


Keywords: Unbiased methods, total-balancing, data adjustment, forest inventory, sampling methods

## 1 Background

For more than half a century, the idea of measuring distance from a random point to the nearest object has been developed. It has often been reviewed in the sampling literature, for instance in books by Pielou (1977), and Bonham (1989). Most of the history of the subject seems to have been developed by ecologists or the mathematicians to whom they brought the problem.

My own interpretation of the method is that it developed roughly as follows:

1) We can see that the average distance to objects, trees for instance, clearly decreases when more objects are added to a fixed tract area - especially if the trees are not extremely clustered. Therefore, distances between random points and objects could be used to estimate the density (meaning objects per unit of land area tree stems in this case).
2) As with many sampling systems, they looked at estimators based on a random distribution, even though this was clearly wrong. Generally, the area around each tree was computed using the distance to the nearest tree $\left(\mathrm{r}_{i}\right)$ by an equation known to be unbiased with a random distribution, then averaged to give area $\mathrm{A}_{t}$. This area around the tree was then used to compute the number of trees in an area as follows:

$$
N=\left(\frac{\text { Tract area }}{A_{t}}\right)
$$

This was highly satisfying for random distributions, although the mathematical proof of such a thing was not easy to follow or explain. Having the equation was enough.
3) A feeling of guilt developed in the ecological circles, since everyone knew that trees and other objects were not randomly distributed. No theoretical approach suggested itself, so a period of simulation followed and examined quite a variety of estimations using the distance $\left(\mathrm{r}_{i}\right)$, such as detailed in Engeman (1994). As in all simulations, it was never "done in our own backyard" so any correction constants could not be trusted - no matter how interesting they might be.

Even with no bias, the method will typically give an answer that is too low. This is because of a high variability when some distances to the tree are very short and therefore give very large individual estimates of N. Although these few very large estimates make the system unbiased, they happen rarely enough that the median answer is typically too low. In this case it is arguably wise to use a biased estimate, which gives a smaller actual error in most cases, and just live with the bias.
4) The problem was extended, in hopes that the variability and any perceived bias would go away. Samplers looked at the $2^{\text {nd }}$-closest tree, the $3^{\text {rd }}$, and generally the " n th closest tree" hoping that the bias would asymptotically go away, and indeed that seemed to be the case.
5) At several times people realized that this was really

[^0]a problem of deducing the area of the average Voronoi polygon around individual trees. Once you had that area, of course, that puts you into the well known realm of Horvitz-Thompson estimators and simplifies everything. A Voronoi polygon is the area around a tree where it is the "closest" tree to any point in the polygon. In fact, the situation could be examined with any shape of polygon around trees, provided that the polygons tessellated the area and you could tell which polygon you fell into with a sample point. Voronoi polygons are simply a very convenient situation to consider.

I have never been able to find a simple procedure for calculating the Voronoi polygon area around a single tree while in the field. Solving the problem for thousands of trees with XY coordinates is easily done and quite efficient by computer algorithms, and you would think that perhaps a simple Excel program must be available to do this in the field using angles and distances to trees. I have not been able to find such a program.

I would suggest that perhaps this is one of those times when we could look at the geometry of the situation and perhaps gain some insight. Before samplers found out that calculus was so impressive to journal editors, they would reason out the geometry of various situations and sometimes came up with some inspired results. Consider Walter Bitterlich's development in the 1940's of Angle-Count Sampling (typically called Variable Plot Sampling) as one example (Bitterlich, 1984). He developed this as a geometry exercise, and it changed forest sampling worldwide. Perhaps this is another example that might benefit from such an approach. Geometric proofs are, after all, a valid type of proof. They are every bit as mathematical as an algebraic or calculus approach, and can be much more illuminating.

## 2 The Geometry

Consider, first, the geometry of selecting a tree linearly "closest" to a random point. Clearly this is a question of falling within a Voronoi polygon in which that tree is "nearest". Where other definitions of "closest" are considered, the geometry remains very similar and the solutions here are basically unchanged.

The average area of such polygons provides the key to estimating the number of trees per hectare. A random point in the area is always located in one and only one of these polygons, and falls within those polygons with probability proportional to their area. Figure 1 illustrates this situation.

## 3 The Problem

The question is: how can we estimate the polygon area by only using a linear distance? If we could detect the distance from the tree to the edge of this polygon,


Figure 1: The "nearest-tree" Voronoi polygon, which is sampled, proportional to its size, by a random point.
a solution becomes fairly simple, and the variability of the estimator is much reduced. Consider the distance $\mathrm{R}_{i}$, which is from the tree to the edge of the polygon. The edge is recognized because it is the point where one or more other trees are the same distance from tree i. A shorter distance $\mathrm{r}_{i}$ has traditionally been used as the distance from a random point to the tree. The larger distance $\mathrm{R}_{i}$ is the distance from a tree (or more generally any fixed point) to the edge of the polygon. This distance has some very fortunate characteristics.

One of the examples in some calculus courses is to establish that the quadratic average $\left(\mathrm{R}_{a}\right)$ of the distances $\mathrm{R}_{i}$ chosen with equal probability from any fixed point (for instance the tree in the polygon) is equal to a circle having radius $\mathrm{R}_{a}$ with exactly the same area as that irregular polygon. This was discussed by Matern (1956), and more recently by Gregoire and Valentine (1995). The polygon does not need to have straight edges for this; but it does in our case, because the edges are formed from bisectors of adjacent trees. For nearest-tree situations it is a very simple polygon with a reference point (the tree) which is easy to identify.

This also leads to the estimate:

$$
\left(R_{a}^{2} \times \pi\right)=\text { polygon area }
$$

If we simply use $\left(R_{i}^{2} \times \pi\right)$ in each case, and then average the areas of these circles, we get an unbiased estimate of $\left(R_{a}^{2} \times \pi\right)$ for polygon area. In other words, we simply treat the distances $\left(\mathrm{R}_{i}\right)$ as circle radii, and average those circle areas. We can use this simple arithmetic average because the angular direction from the tree to the polygon edge was randomly chosen with equal probability.


Figure 2: A circle having a radius $\left(\mathrm{R}_{a}\right)$ equivalent to the quadratic average of all possible distances $\sqrt{\frac{\sum R_{i}^{2}}{n}}$, has an area equal to the area of that irregular polygon.

If the ray outward from the tree was not randomly chosen (such as when it was chosen by going through a random point) we would have to weight the individual distances to compute the same expected value. Here again, we have only to refer to previous work. Walter Bitterlich taught foresters how to select circles proportional to their area and how to use the results. This simple geometry problem was solved by using an angle gauge to choose trees at a random point. A random point chooses the larger circles (radii) by the square of the radius involved.

If we wanted to have the arithmetic average of the radii as if the radii were chosen equally, the first suggestion for this seems to have come from Hirata (1956). We simply take the harmonic mean of the squared radii, because the weighting of their selection was made with probability proportional to the squared distance. It is easy to imagine the weight being proportional to a small wedge extending from the tree outwards, so the probability of a point falling into this area is proportional to the square of the distance:

$$
R_{a}=\sqrt{\frac{1}{\left(\frac{\sum\left[\frac{1}{R_{i}^{2}}\right]}{n}\right)}}
$$

This is the unbiased estimate of the arithmetic mean of equally chosen radii, even though the radii used in this computation were chosen proportional to their squared length by going from the tree through a randomly chosen
point in the polygon.
How could we do this in the field? Our problem is simply to sample for the average circle area $\left(R_{a}^{2} \times \pi\right)$ using distances from the tree to the polygon edge. One way to do this is:

1) Select a random point and go to the nearest tree.
2) From the tree, select a random angle, and go in that direction until the edge of the polygon is encountered. This is the first point where another tree would be equally far away $\left(\mathrm{R}_{i}\right)$.
This "random direction" step can be skipped if you use the harmonic mean just described, in which case the distance $\mathrm{R}_{i}$ is from the tree through the sample point to the edge of the polygon. This simplifies field work.
3) Measure $R_{i}$, as an estimate of a circle radius equal to the polygon area. The average of these squared radii (weighted harmonically, if necessary) is $\mathrm{R}_{a}^{2}$.
( $R_{a}^{2} \times \pi$ ) then estimates the average polygon area around individual trees.
4) From this, the number of trees/ha can be calculated.

Other estimates of volumes, values and other characteristics are similarly best imagined geometrically, but will be more fully described in future papers. To those who are familiar with Variable Plot sampling, these are easily imagined as Volume to Basal Area Ratios (VBARs). When averaged, these can simply be multiplied by stand area in order to produce totals for the tract.

It is relatively easy to do such calculations. The main deviation from previous work comes from viewing the problem as a sample of various size circles, rather than using any assumption at all about tree distribution. Note that there is absolutely no restriction at all on the distribution of trees.

At this point, we have the same form of equation as has always existed for point to plant areas and numbers per hectare. The only difference is that the circle area derived from point to the plant distances $\left(\mathrm{r}_{i}\right)$ was doubled. This was used because when we choose a random point in a circle, the average area of $\left(r_{i}^{2} \times \pi\right)$ is $1 / 2$ of the area $\left(R_{i}^{2} \times \pi\right)$.

The problem with this historical estimator is that it is quite variable. The distance to objects can obviously be very small, and when dealing with reciprocal squares this caused high variability. There was a way to solve this problem, and it involved going beyond the nearest tree, which might be quite close, and going to the $2^{\text {nd }}$, $3^{d}$ or more generally the " n th nearest-tree".

This problem is also best viewed as a geometry problem. Although some have apparently viewed this distance to the $\mathrm{n}^{\text {th }}$ tree as a kind of "plot radius" (Lynch, 2003), I believe that this does not provide insight into the actual geometry. Lately, several authors, for instance Magnussen (2008) and Kleinn (2006) deduced that this involved "order-k" Voronoi polygons (Okabe, et al, 1999, chapter 3, page 152) around trees, but admitted that calculating these in the field was even more impractical. Indeed, measuring the polygon is too awkward to contemplate, but sampling for it is not. Using many trees in a larger polygon, rather than just one $\mathrm{n}^{t h}$-nearest tree certainly complicates imagining the geometry.

If you consider the Voronoi polygons around the nearest tree, and ask what is the polygon where this tree is the " $2{ }^{n d}$-nearest-tree", the geometry begins to clear up. In this illustration, I have taken the polygons for trees bordering an example tree and calculated the parts of these where the example tree is the $2^{n d}$-nearest-tree. This can be done by hand, and I am sure that it could be done quickly and more accurately by a GIS system, which would be especially necessary for larger " n th nearest" situations. The analysis depends upon individually eliminating trees, then dividing that tree's polygon among other trees. This process adds what I will call "slivers" along the edge each of the original Voronoi polygon. It is within these slivers that the tree is the $2^{\text {nd }}$-nearest-tree. You do not need to know this area or the boundaries, because you know when you fall into that polygon (because that example tree is the second closest), but visualizing the geometry reveals a lot about why it works well.

The graph from this process produces a "halo" (Fig. 3) of slivers which surrounds each tree. Here they are illustrated around just one of the trees.

The consequence of this is that the starting point for the distance to the " $n$-th nearest-tree" must lie within these slivers. The outer border of the halo forms a new polygon consisting of the inner original polygon plus the added slivers. On the average, these larger polygons are exactly twice the size of the inner polygons describing the nearest trees. The smaller slivers add up to exactly the tract area, and are all allocated to one and only one tree. The interior parts of the original "nearesttree" polygon would be divided into slivers which would select some other tree as the $2^{n d}$-closest. Therefore, the original polygons plus the sliver areas that border them amount to twice the area of the tract, and with the same number of trees those polygons have an average exactly twice as large.

We therefore have the same solution as before. If we measure the radius to the edge of this larger polygon, then calculate the average area, we will estimate twice the area of an average nearest-tree polygon. The same


Figure 3: A "halo" of slivers forms along the border of the original polygon to indicate where it would be chosen as the "second-closest" tree.
process is used, but the area is just divided by two before you calculate numbers of trees. The same reasoning, of course, applies to the $3^{\text {rd }}, 4^{\text {th }}$ or $\mathrm{n}^{\text {th }}$ closest tree. The halos of slivers get thinner, and occur at greater distances from the tree. The slivers tessellate the area as if they were a large stained-glass window with interlacing halos of different colors, each assigned to different trees.

What we would prefer is the distance from the tree to the edge of this larger polygon $\left(\mathrm{R}_{i}\right)$, but the simple distance from the random point to the tree $\left(r_{i}\right)$ is at least restricted by the width of the slivers along the border of the polygon. This shorter distance, if used directly, would lead to an estimate of an average polygon which is too small, and therefore would estimate too large a number of trees. Although any bias from using $\mathrm{r}_{i}$ rather than $\mathrm{R}_{i}$ may be smaller, and although it reduces as we go to the $4^{t h}, 5^{t h}, 6^{t h}$ tree and so on, we would prefer the distance $\mathrm{R}_{i}$ because it is unbiased. To find the actual polygon edge of the larger polygon we should back away from the tree until it ties with another tree as the " n th closest tree".

The bias caused by using a shorter distance ( $\mathrm{r}_{i}$ ) has caused some to suggest that an additional distance be added to each measurement, which can reduce the bias. This was usually visualized as using a slightly larger "fixed plot" with the n trees inside it, since the distance barely includes the $\mathrm{n}^{\text {th }}$ tree. I do not think that this view is useful for understanding the process, but some adjustment would clearly help to reduce the bias.

When using the $\mathrm{n}^{\text {th }}$ closest tree approach the variabil-
ity has been reduced, and at some point the bias becomes negligible because these slivers are too slim to create a great deal of difference in the distance to the sample point versus the correct distance to the polygon edge. It is a classic tradeoff, an unbiased method that is more awkward in the field compared to a biased estimate that is relatively stable and has simple field measurements.

I must admit to being one who would use the biased method. On the other hand, what would happen if we had a simple instrument or method that would tell us when we crossed that invisible boundary where the tree went from the $\mathrm{n}^{\text {th }}$ nearest to the $(\mathrm{n}+1)^{\text {th }}$ nearest? We would then have an unbiased system with desirable variability characteristics. All we need to be aware of this possibility is to view the geometry in such a way as to see the actual situation. Bitterlich found a way to tell when he was inside an invisible circle that was a multiple of the stem area without distance measurements or calculations, by simply using an angle to view the tree. When we look at the nearest-tree process as a geometry exercise, perhaps someone else will show similar ingenuity. There are obvious extensions of this geometric view to other items besides simple tree numbers. I think that this view is general, useful, and puts the mathematics into context in a way that pure mathematical approaches do not.

It was a large breakthrough when the scientific community discovered the concept of analytic geometry. Have we forgotten the geometry part of that insight? I think that perhaps we have. The reason that this problem has essentially gone unsolved for so very long is that it does not yield readily to a purely mathematical solution without the geometrical insight. Variable Plot sampling was an enormous breakthrough in forest sampling. I believe that this was because it was essentially a geometrical problem solved by a geometrical insight. I think the nearest-neighbor problem is the same, and that there are still many problems like these.

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