MACHINE LEARNING 101

Rev#	Date	Revised by	Comments
1.0	14/Feb/2019	Khelil Sator	Initial version of the doc Excel file to compute three-sigma rule Excel file to compute one iteration of k-means with k=2 Machine learning demos for anomaly detection using Healthbot https://github.com/ksator/junos monitoring with healthbot/wiki/Machine-learning
1.1	15/Feb/2019	Khelil Sator	Added anomaly detection (HB 1.0) content from Vijay Gadde
1.2	14/April/2019	Khelil Sator	Added outlier detection details (available with HealthBot 2.0) - k-Fold Cross-Validation - K-fold Three-sigma - DBSCAN (Density-Based Spatial Clustering of Applications with Noise)

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Why this doc

Using HealthBot, it is easy to collect data from the network devices, to store the data collected in a database, and to use machine learning for anomaly detection.

The purpose of this doc is to help peoples with no machine learning background to better understand machine learning basics.

It may seem weird to describe how to calculate a standard deviation, a variation, a Euclidean distance, an argmin, ... but this is required to fully understand how k-means clustering works.

Anomaly detection vs outlier detection

HelthBot supports machine learnings for anomaly detection and for outlier detection.

HealthBot supports the following machine learning algorithms for anomaly detection:

- Three-sigma rule
- k-means for anomaly detection

HealthBot supports the following machine learning algorithms for outlier detection:

- DBSCAN (Density-Based Spatial Clustering of Applications with Noise)
- K-fold Three-sigma ("K-Fold Cross-Validation" using "Three-sigma")

Anomaly detection and outlier detection are both about detecting anomalies.

In HealthBot terminology:

- anomaly detection is time based. It compares new data points from a device vs data points collected from the same device during a learning period.
- outlier detection is group based. It analyzes data from a device during a learning Period vs data from other devices during the same learning period

Supervised learning

Learning with a teacher.

The machine learning algorithm learns on a labeled dataset

Unsupervised learning

Learning without a teacher.

The machine learning uses unlabeled dataset.

The advantage of using an unsupervised technique is that we do not need to have labeled data, i.e., we do not need to create a training dataset that contains examples of outliers.

k-means clustering and DBSCAN are unsupervised clustering machine learning algorithms.

They group the data that has not been previously labelled, classified or categorized.

Machine learning model

This is the output generated when you train your machine learning algorithm with your training dataset.

The machine learning model is what you get when you run the machine learning algorithm over your training data.

Once The machine learning model is built, it can be used to classify new data points.

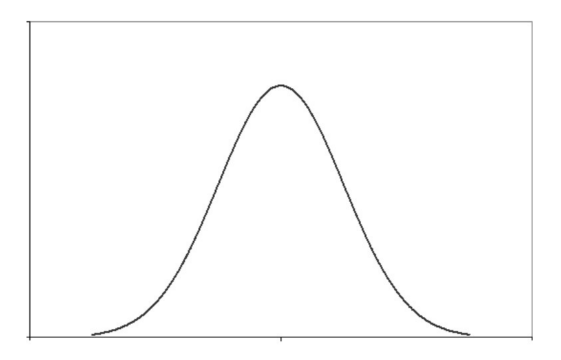
Normal distribution

Also called: Bell curve, Gaussian distribution, Gauss distribution, Laplace-Gauss distribution

A "normally distributed" data set has most of the data aggregates around its mean in a symmetric fashion. Values become less and less likely to occur the farther they are from the mean.

Think about a factory producing 1 kg bags of sugar. They won't always make each exactly 1 kg. In reality, the bags are around 1 kg. Most of the time they will be very close to 1 kg, and very rarely far from 1 Kg.

Example of normally distributed data: height of adults



Standard deviation

Overview

A measure that is used to quantify the amount of variation of a set of data values.

A low standard deviation indicates that the data points tend to be close to the mean of the set.

A high standard deviation indicates that the data points are spread out over a wider range of values.

Its symbol is σ (the greek letter sigma)

The formula for standard deviation (SD) is

$$ext{SD} = \sqrt{rac{\sum |x-\mu|^2}{N}}$$

where \sum means "sum of", x is a value in the data set, μ is the mean of the data set, and N is the number of data points in the population.

Calculate it by hand

Data set = 101, 102, 106, 107

```
Mean = (101 + 102 + 106 + 107)/4 = 104

((101-104)^2 + (102-104)^2 + (106-104)^2 + (107-104)^2)/4 = (9 + 4 + 4 + 9)/4 = 6.5

standard deviation = \sqrt{6.5} = 2.549
```

Variance

Overview

As indicated above, the Standard Deviation is a measure of how spread out numbers are. Think about the average difference around the mean.

The standard deviation is the square root of the variance. As example: if SD = 3, variance = 9

The variance is the average of the squared differences from the Mean.

Calculate it by hand

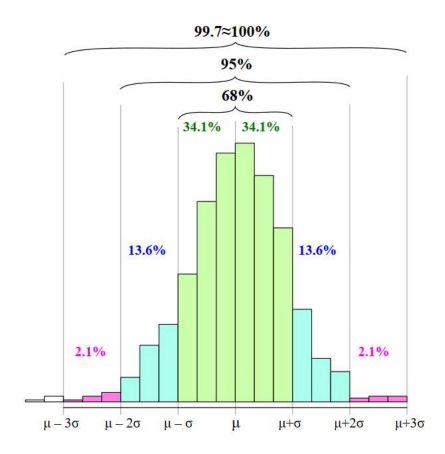
Data set = 101, 102, 106, 107

Mean = (101+102+106+107)/4=104Var = $((101-104)^2 + (102-104)^2 + (106-104)^2 + (107-104)^2)/4 = (9+4+4+9)/4 = 6.5$

68-95-99.7 rule

With a normal data set (normally distributed) (as example: height of adults):

- 68.27% of the values of the data set are in a band of two standard deviations around the mean (mean 1 standard deviation <-> mean + 1 standard deviation)
- 95% of the values of the data set are in a band of four standard deviations around the mean (mean 2 standard deviations <-> mean + 2 standard deviations)
- 99.7% of the values of the data set are in a band of six standard deviations around the mean (mean 3 standard deviations <-> mean + 3 standard deviations)



three-sigma rule

Overview

sigma is the greek letter σ . This is also the Standard Deviation symbol

HealthBot uses 3-sigma rule for anomaly detection.

Three-sigma rule classifies a new data point as "normal" if it is between the (mean -3 * standard deviations) and (mean +3 * standard deviations)

Three-sigma rule classifies a new data point as "abnormal" if it is outside this range.

mean: mean of the data set

SD: standard deviation of the data set

abs: absolute value

x: a new data point

If abs(x - mean) > (3 * SD) then tree-sigma classifies x as abnormal

If abs(x - mean) < (3 * SD) then tree-sigma classifies x as normal

Calculate it by hand

Data points = 101, 102, 106, 107

Mean = (101 +102 + 106 + 107)/4=104

 $((101-104)^2 + (102-104)^2 + (106-104)^2 + (107-104)^2)/4 = (9+4+4+9)/4 = 6.5$

standard deviation = $\sqrt{6.5}$ = 2.54

3 * standard deviation = 7.62

mean – 3 * standard deviation = 96.38

mean + 3 * standard deviation = 111.62

Three-sigma rule classifies a new data point as "normal" if it is between 96.38 and 111.62

Three-sigma rule classifies a new data point as "abnormal" if it is outside this range.

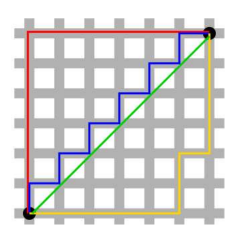
HealthBot rule example

HealthBot rule using OpenConfig telemetry to monitor the number of BGP prefixes received per peer, and using a dynamic threshold computed by three-sigma to detect anomaly: https://github.com/ksator/junos monitoring with healthbot/blob/master/rules/check-bgp-routes-

with-3-sigma.rule

Euclidean distance vs Manhattan distance

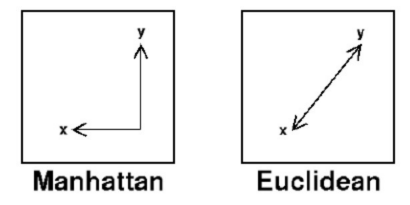
Overview



In green: Euclidean distance between X and Y. This is the shortest path between X and Y. crow flies.

In Red, Yellow, Blue: Manhattan distance between X and Y. This is the shortest path a car could take between X and Y in Manhattan

Note: Red, Yellow, Blue paths have the same length.



Calculate Euclidean distance by hand

We generally use a "double vertical line" notation for Euclidean distance. So the Euclidean distance between X and Y is d(X,Y) = ||Y - X||

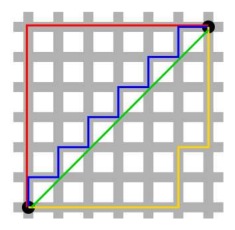
Euclidean distance formula:

$$\sqrt{\sum_{i=1}^n (x_i-y_i)^2}$$

If X and Y are defined with 2 variables, let's say $X = (X_1, X_2)$ and $Y = (Y_1, Y_2)$, the Euclidean distance between X and Y is equal to: $\sqrt{((X_1-Y_1)^2+(X_2-Y_2)^2)}$

Just remember Pythagorean theorem! That's it.

If X = (0,0) and Y = (6,6), the Euclidean distance between X and Y is equal to $\sqrt{(36+36)}$ = 8.36 See the green path to better understand the above example



Calculate Manhattan distance by hand

Manhattan distance formula:

If X and Y have n dimensions:

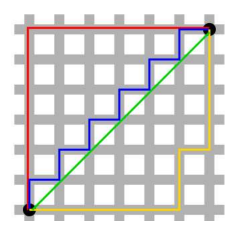
$$\sum_{i=1}^n |x_i-y_i|$$

If X and Y are defined with 2 variables, let's say $X = (X_1, X_2)$ and $Y = (Y_1, Y_2)$, the Manhattan distance between X and Y is equal to:

$$|Y_1 - X_1| + |Y_2 - X_2|$$

If X = (0,0) and Y = (6,6), the Manhattan distance between X and Y is equal to 12

See the red/blue/yellow paths to better understand the above example.



argmax

Also called arg max.

```
\operatorname{arg\,max}_{x} f(x)
```

In mathematics, this is the inputs, or arguments, at which the output of the function f is maximized example:

```
f(x) = -|x|
```

f(x) value is maximized with x = 0

argmin

Also called arg min.

```
arg min_x f(x)
```

This is the value of x for which f(x) attains it's minimum.

example:

f(x) = 2 * x

f(x) value is minimized with x = 0

"k-means clustering" vs "k-means for anomaly detection"

"k-means clustering" and "k-means for anomaly detection" are two different things.

"k-means clustering" splits n data points into k clusters.

"k-means for anomaly detection" uses "k-means clustering" and others building blocks as well to identify "abnormal" data points (data points significantly different from the majority of the data).

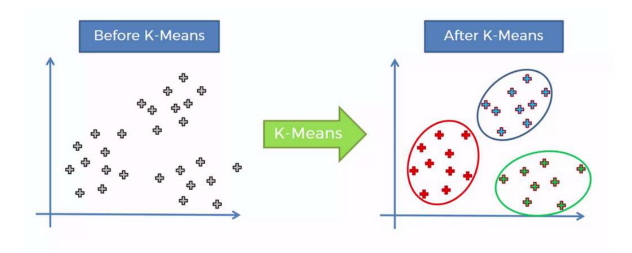
HealthBot uses "k-means for anomaly detection" to build a model and to classify new data points as normal or abnormal.

k-means clustering

Overview

k-means clustering splits N data points into K groups (called clusters).

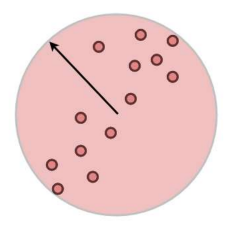
You need to specify how many clusters you want.



A cluster is a group of data points.

Each cluster has a center, called the centroid. A cluster centroid is the mean of a cluster (average across all the data points in the cluster).

The radius of a cluster is the maximum distance between all the points and the centroid.



Distance between clusters = distance between centroids.

k-means clustering is more difficult than three-sigma rule. k-means looks difficult, but it is not so difficult. It uses a basic iterative process. We will run it later on in this doc.

k-means clustering splits N data points into K clusters. Each data point will belong to a cluster. This is based on the nearest centroid. This uses Euclidean distance.

The objective is to find the most compact partitioning of the data set into k partitions. k-means makes compacts clusters. It minimizes the radius of clusters.

The objective is to minimize the variance within each cluster.

Clusters are well separated from each other. It maximizes the average inter-cluster distance.

Formula

Given a set of data points $(\mathbf{x}_1, \mathbf{x}_2, ..., \mathbf{x}_n)$, k-means partitions the n data points into k $(\leq n)$ clusters $\mathbf{S} = \{S_1, S_2, ..., S_k\}$ while minimizing variance within each cluster (i.e while minimizing the Euclidean distance between each data point and the centroid of the cluster it belongs to)

The objective is to find:

$$\operatorname*{arg\,min}_{\mathbf{S}} \sum_{i=1}^k \sum_{\mathbf{x}_j \in S_i} \|\mathbf{x}_j - \boldsymbol{\mu}_i\|^2$$

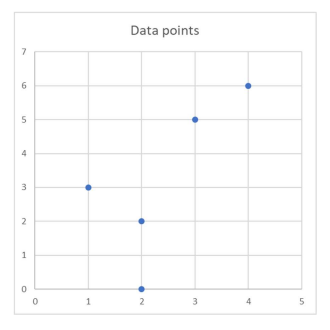
where μ_i is the mean of points in S_i .

Calculate it by hand

As indicated, k-means looks difficult, but it is not so difficult. It uses a basic iterative process. Let's run it. let's run k-means, with k=2, with the below data points.

In this example each data point is defined with 2 variables.

Data points	Variable 1	Variable 2
D1	2	0
D2	1	3
D3	3	5
D4	2	2
D5	4	6



So:

we have 5 data points (D1, D2, D3, D4, D5).

We want 2 clusters (cluster 1 and cluster 2).

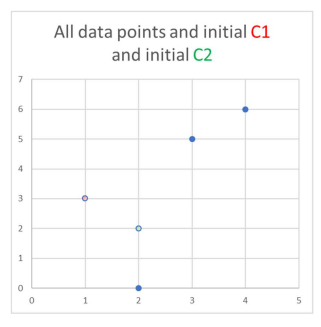
We will calculate the centroid for each of the 2 clusters. Let's call the centroids C1 and C2. C1 is the centroid for Cluster 1 and C2 is the centroid for Cluster 2.

For each cluster we will find out the centroid and which data points belongs to the cluster.

Let's start:

k-means generates randomly 2 initial centroids (or select randomly 2 existing data points as the initial centroids).

In this demo, we will select 2 existing data points (D2 and D4) as the initial centroids. Let's say initial C1 = D2 and initial C2 = D4 so initial C1 is (1,3) and initial C2 is (2,2)



Iteration 1

Let's calculate the Euclidean Distance between each data points and the current centroids.

Current centroids are: C1 (1,3) and C2 (2,2)

Details for d1 and d2: $\sqrt{((2-1)^2+(3-0)^2)} = \sqrt{((1)^2+(3)^2)} = \sqrt{(1+9)} = \sqrt{10} = 3.16227766$

Details for d1 and d4: $\sqrt{((2-2)^2+(2-0)^2)} = \sqrt{((0)^2+(2)^2)} = \sqrt{(0+4)} = \sqrt{4} = 2$

After calculating the Euclidean distance between all data points each data points and the current centroids, we get:

Data points	Euclidean distance to current C1	Euclidean distance to current C2
D1	3.16227766	2
D2	0	1.41421356
D3	2.82842712	3.16227766
D4	1.41421356	0
D5	4.24264069	4.47213595

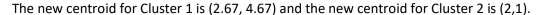
Let's group the data points in clusters, based on the closest centroid (Euclidean distance)

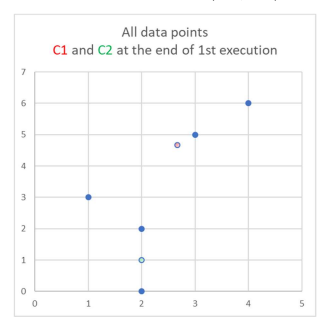
Cluster 1: D2, D3, D5.

Cluster 2: D1, D4

For each cluster, let's calculate the mean for Variable 1 and Variable 2. This will be the new centroids.

Cluster	Data points	Data points Mean (Variable 1)	Data points Mean (Variable 2)
Cluster 1	D2, D3, D5	2.67	4.67
Cluster 2	D1, D4	2	1





1st iteration is done.

Let's repeat these steps until either k-means converges (i.e new centroids don't change anymore) or until we hit the max number of iterations.

2nd iteration.

Let's calculate the Euclidean Distance between each data points and the current centroids.

Current centroids are: C1 (2.67, 4.67) and C2 (2,1).

Data points	Euclidean distance to current C1	Euclidean distance to current C2
D1	4.71781729	1
D2	2.36173665	2.23606798
D3	0.46669048	4.12310563
D4	2.75278041	1
D5	1.88090404	5.38516481

Let's group the data points in clusters, based on the closest centroid (Euclidean distance)

Cluster 1: D3, D5

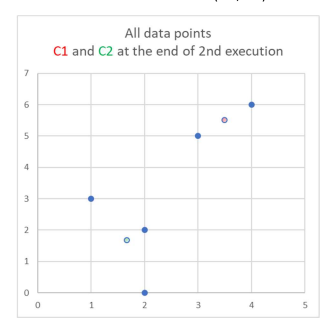
Cluster 2: D1, D2, D4

For each cluster, let's calculate the mean for Variable 1 and Variable 2. This will be the new centroids.

Cluster	Data points	Data points Mean (Variable 1)	Data points Mean (Variable 2)
Cluster 1	D3, D5	3.5	5.5

Cluster 2	D1, D2, D4	1.67	1.67
Cidotei 2	01,02,01	07	1.07

The new centroid for Cluster 1 is (3.5, 5.5) and the new centroid for Cluster 2 is (1.67,1.67).



So the centroids changed, so the algorithm did not yet converged.

2nd iteration is done.

Let's repeat these steps until either k-means converge (i.e new centroids don't change) or we hit the max number of iterations.

3rd iteration:

Current centroids are: C1 (3.5, 5.5) and C2 (1.67,1.67)

Let's calculate the Euclidean Distance between each data points and the current centroids.

Data points	Euclidean distance to current C1	Euclidean distance to current C2
D1	5.70087713	1.70229257
D2	3.53553391	1.48922799
D3	0.70710678	3.58577746
D4	3.80788655	0.46669048
D5	0.70710678	4.91709264

Let's group the data points in clusters, based on the closest centroid (Euclidean distance)

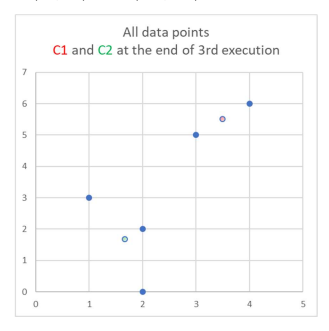
Cluster 2: D3, D5

Cluster 1: D1, D2, D4

For each cluster, let's calculate the mean for Variable 1 and Variable 2. This will be the new centroids.

Cluster	Data points	Data points Mean (Variable 1)	Data points Mean (Variable 2)
Cluster 1	D3, D5	3.5	5.5
Cluster 2	D1, D2, D4	1.67	1.67

C1 (3.5, 5.5) and C2 (1.67,1.67)



C1 and C2 did not change with this new iteration!

K-means converged, in 3 iterations.

Done!

We have our 2 clusters: Cluster 1 and cluster 2

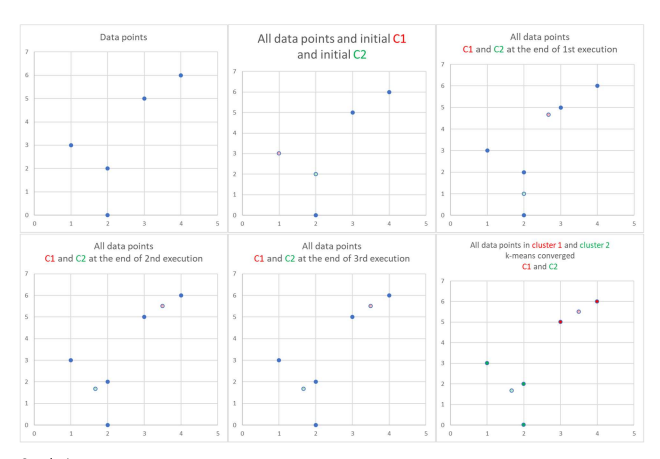
We have our 2 Centroids C1(3.5, 5.5) and C2(1.67, 1.67)

D3, D5 are in cluster 1.

D1, D2, D4 are in Cluster 2.



Quick visual recap:



Conclusion:

When a new data point will be added (D6) to the data set, k-means will classify it in cluster 1 or in cluster 2 based on the shortest Euclidean distance between D6 and C1 vs D6 and C2.

k-means for anomaly detection

Overview

"k-means for anomaly detection" uses "k-means clustering" and others building blocks as well to identify "abnormal" data points (data points significantly different from the majority of the data).

HealthBot uses "k-means for anomaly detection"

HealthBot rule example

Healthbot uses "k-means for anomaly detection".

HealthBot rule using OpenConfig telemetry to monitor the number of BGP prefixes received per peer, and using a dynamic threshold computed by k-means to detect anomaly: https://github.com/ksator/junos_monitoring_with_healthbot/blob/master/rules/check-bgp-routes-with-k-means.rule

k-Fold Cross-Validation (CV)

CV uses a single parameter called k.

It works like this:

it splits the dataset into k groups.

For each unique group:

- Take the group as a test data set
- Take the remaining groups as a training data set
- Use the on the training set to build the model, and then use the test set and evaluate

Example:

A dataset 6 datapoints: [0.1, 0.2, 0.3, 0.4, 0.5, 0.6]

The first step is to pick a value for k in order to determine the number of folds used to split the dataset.

Here, we will use a value of k=3. so we split the dataset into 3 groups. each group will have an equal number of 2 observations.

For example:

Fold1: [0.5, 0.2]

Fold2: [0.1, 0.3]

Fold3: [0.4, 0.6]

Three models are built and evaluated.

Model1: Trained on Fold1 + Fold2, Tested on Fold3

Model2: Trained on Fold2 + Fold3, Tested on Fold1

Model3: Trained on Fold1 + Fold3, Tested on Fold2

K-fold Three-sigma

"K-fold Three-sigma" is "k-Fold Cross-Validation" using "Three-sigma"

The algorithm used by CV is "tree-sigma" rule.

HealthBot can use "K-fold Three-sigma" for outlier detection

We need to indicate to HealthBot which dataset to use (as example "number of BGP prefixes sent" or "cpu load").

For this dataset, "K-fold Three-sigma" analyzes data from a device during a configurable learning Period vs data from other devices during the same learning period

K is the number of devices: if the group has 4 devices, then k=4, so HealthBot will use "4-fold three-sigma".

In that case, 4 models are built and evaluated by HealthBot.

Model1: Trained with the data points collected during the learning period from device 1 and device 2 and device 3, then tested with the data points collected during the learning period from device 4

Model2: Trained with the data points collected during the learning period from device 1 and device 2 and device 4, then tested with the data points collected during the learning period from device 3

Model3: Trained with the data points collected during the learning period from device 1 and device 3 and device 4, then tested with the data points collected during the learning period from device 2

Model4: Trained with the data points collected during the learning period from device 2 and device 3 and device 4, then tested with the data points collected during the learning period from device 1

HealthBot returns 1 if it detects an outlier, and 0 if there is no outlier detected.

DBSCAN (Density-Based Spatial Clustering of Applications with Noise)

It is an unsupervised machine learning algorithm.

It is a density-based clustering algorithm.

It groups datapoints that are in regions with many nearby neighbors.

It groups datapoints in such a way that datapoints in the same cluster are more similar to each other than those in other clusters.

Clusters are dense groups of points. Clusters are dense regions in the data space, separated by regions of lower density

If a point belongs to a cluster, it should be near to lots of other points in that cluster.

It marks datapoints in lower density regions as outliers.

It uses time series data and detect outliers. We collect a window of data from a TSDB. This window is then passed to the DBSCAN algorithm, which returns the set of datapoints considered outliers.

It works like this:

First, we choose two parameters, a number <u>epsilon</u> (distance) and a number <u>minPoints</u> (minimum cluster size). epsilon is a letter of the Greek alphabet.

We then begin by picking an arbitrary point in our dataset.

If there are at least minPoints datapoints within a distance of epsilon from this datapoint, this is a high density region and a cluster is formed. i.e if there are more than minPoints points within a distance of epsilon from that point (including the original point itself), we consider all of them to be part of a "cluster".

We then expand that cluster by checking all of the new points and seeing if they too have more than minPoints points within a distance of epsilon, growing the cluster recursively if so.

Eventually, we run out of points to add to the cluster. We then pick a new arbitrary point and repeat the process.

Now, it's entirely possible that a point we pick has fewer than minPoints points in its epsilon range, and is also not a part of any other cluster: in that case, it's considered a "noise point" (outlier) not belonging to any cluster.

epsilon and minPoints remain the same while the algorithm is running.

For more information about this algorithm you can read

https://www.naftaliharris.com/blog/visualizing-dbscan-clustering/

https://medium.com/netflix-techblog/tracking-down-the-villains-outlier-detection-at-netflix-40360b31732

Machine learning usage with HealthBot

Overview

HelthBot supports machine learnings for anomaly detection and for outlier detection.

HealthBot supports the following machine learning algorithms for anomaly detection:

- Three-sigma rule
- k-means for anomaly detection

HealthBot supports the following machine learning algorithms for outlier detection:

- DBSCAN (Density-Based Spatial Clustering of Applications with Noise)
- K-fold Three-sigma ("K-Fold Cross-Validation" using "Three-sigma")

Anomaly detection and outlier detection are both about detecting anomalies.

In HealthBot terminology:

- anomaly detection is time based. It compares new data points from a device vs data points collected from the same device during a learning period.
- outlier detection is group based. It analyzes data from a device during a learning Period vs data from other devices during the same learning period

How to use machine learning for anomaly detection with HealthBot

HealthBot can use machine learning for anomaly detection (to classify new data points as `normal` or `abnormal`).

HealthBot supports the following machine learning algorithms:

- Three-sigma for anomaly detection
- k-means for anomaly detection.

For each algorithm, we need to configure:

- the learning-period
- the pattern-periodicity

Machine learning model:

- This is the output generated when you train your machine learning algorithm with your training data-set. The machine learning model is what you get when you run the machine learning algorithm over your training data.
- Training the machine learning algorithm to build models is done every day by HealthBot at midnight.
- The machine learning model is then used in production to handle new data points.

Learning-period is configured to determine how much historical data the machine learning algorithm uses to build the models.

For example, if learning period is 7 days, every time we build the models, we fetch past 7 days of data and we train the machine learning algorithm to build new models.

if the learning period is 7 days, when learning is triggered the 12th Feb 2019 (00:00) to build new models, we fetch data from 5th Feb 2019 00:00 to 12th Feb 2019 00:00 and train the machine learning algorithm.

If the learning period is 1 month, when learning is triggered the 12th Feb 2019 (00:00), we fetch data from 12th Jan 2019 00:00 to 12th Feb 2019 00:00 and train the machine learning algorithm to build new models.

pattern-periodicity:

- If we configure '1D' 'pattern-periodicity', it means that data of each day of the week has different patterns. So HealthBot creates 7 buckets, one for each day in a week (Monday, Tuesday, ...).
- If we configure '1H' 'pattern-periodicity', it means that regardless of which day, week or month, data of every hour has specific pattern. So HealthBot creates 24 buckets, one for each hour (00:00-00:59, 1:00-1:59, 2:00-2:59 ... 23:00-23:59).
- If we configure '1D 1H' 'pattern-periodicity', it means that for every day, data for each hour has different pattern. So, we would have 7 * 24 = 168 buckets. For Monday 24 buckets (1 for every hour), for Tuesday 24 buckets (1 for every hour) and so on. Here it does not matter in which months we are collecting the data from.
- ...

For a machine learning algorithm to be able to build a model for a bucket, it requires a minimum number of data points. This depends on what algorithm we are using.

- 3-sigma requires only a couple of points per bucket.
- k-mean requires at least 32 data points per bucket.

Once the models are built, they are used in production to handle new data points. HealthBot uses machine learning for anomaly detection with dynamic thresholds. HealthBot uses the models it built to classify new data points as `normal` or `abnormal`.

If the data set used to build the models was small, the result will not be accurate. With larger data set to build the models, results accuracy will increase.

HealthBot builds models based on pattern periodicity. With a pattern periodicity of 1 hour, 24 buckets are formed (one for each hour).

Learning period is only used to fetch the data to build the models. For example, if we have data for past one year, but we have configured learning period as 30 days, in this case, though we have data for past one year, at every midnight when we are re-building the models, we will fetch only past 30 days of data, bucketize the data points based on the pattern periodicity and build one model for each bucket.

Minimum learning period can be the time where you can have minimum data points required to build a model for each bucket. But keep learning period larger to have more accurate results.

Let's take an example:

Let's say at 13:00 p.m we configure HealthBot to collect data from network devices every 5 seconds and to use k-means for anomaly detection with dynamic thresholds with a pattern periodicity of 1 hour.

24 buckets are formed (one for each hour).

Models are built daily at midnight. Until then HealthBot will return -1 as there is no model built yet for any of the buckets. So, during the first day, HealthBot will return -1 for each new data point.

At midnight the first day, k-means will try to build the models:

- The models for hours 00 12 can't be built yet as there is no data collected yet to build these models. So, during the second day, HealthBot will keep returning -1 until 12:59 p.m.
- Since k-means will have enough data per bucket/hour for hours 13 23 (k-means requires at least 32 data points per bucket, and we have 12 * 60 = 720 data points for each of these buckets), it will build the models for these 11 hours/buckets only. From 13:00 p.m the second day, since the models are available, for each new data point, HealthBot will return 0 (normal) or 1 (abnormal).

So, the second day, HealthBot will return -1 for hours 00 – 12 and 0 and/or 1 for hours 13 - 23

At midnight the second day, we have data for all hours 0-23 (one day of data for hours 0-12 hours and 2 days of data for 13-23 hours).

Now since we have enough data points for all the hours/buckets, models for all the hours will be built, and HealthBot will return 0 (normal) or 1 (abnormal) for each new data point it will receive.

Examples of machine learning for anomaly detection with HealthBot

HealthBot rule using OpenConfig telemetry to monitor BGP sessions state (without machine learning to detect anomaly)

https://github.com/ksator/junos_monitoring_with_healthbot/blob/master/rules/check-bgp-state-using-openconfig.rule

HealthBot rule using OpenConfig telemetry to monitor the number of BGP prefixes received per peer, and using a static threshold (provided as a variable) without machine learning to detect anomaly: https://github.com/ksator/junos monitoring with healthbot/blob/master/rules/check-bgp-routes.rule

HealthBot rule using OpenConfig telemetry to monitor the number of BGP prefixes received per peer, and using a dynamic threshold computed by three-sigma to detect anomaly:

https://github.com/ksator/junos_monitoring_with_healthbot/blob/master/rules/check-bgp-routes-with-3-sigma.rule

HealthBot rule using OpenConfig telemetry to monitor the number of BGP prefixes received per peer, and using a dynamic threshold computed by k-means to detect anomaly:

https://github.com/ksator/junos_monitoring_with_healthbot/blob/master/rules/check-bgp-routes-with-k-means.rule

HealthBot playbook using the above 4 rules:

https://github.com/ksator/junos_monitoring_with_healthbot/blob/master/playbooks/machine-learning-for-bgp.playbook

This link provides more details about above rules and playbook. It includes also a script to update continuously the number of BGP prefixes in your network

https://github.com/ksator/junos_monitoring_with_healthbot/wiki/Machine-learning-for-anomaly-detection-(demo-with-BGP-prefixes-received)