

## Lab1: Pima Indians Diabetes Database

This lab uses the classic [Pima Indians Diabetes dataset](#), which contains medical measurements collected from female patients of Pima Indian heritage.

### Dataset description

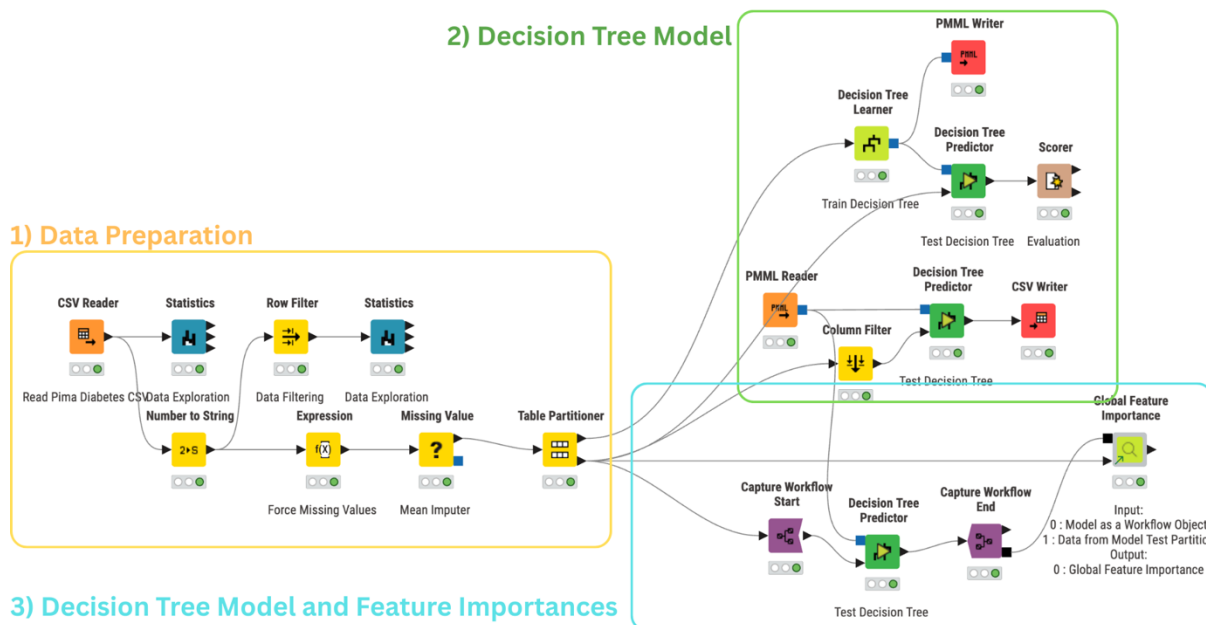
The Pima Indians Diabetes dataset contains 768 records, each representing one female patient of Pima Indian heritage aged 21 years or older. For each patient, the following variables are recorded:

- **Pregnancies**  
Number of times the patient has been pregnant (integer count).
- **Glucose**  
2-hour plasma glucose concentration (mg/dL) from an oral glucose tolerance test.
- **BloodPressure**  
Diastolic blood pressure (mm Hg).
- **SkinThickness**  
Triceps skinfold thickness (mm), a proxy for body fat.
- **Insulin**  
2-hour serum insulin ( $\mu$ U/mL).
- **BMI**  
Body Mass Index, defined as weight in kg divided by the square of height in meters ( $\text{kg}/\text{m}^2$ ).
- **DiabetesPedigreeFunction**  
A derived score that estimates the patient's hereditary risk of diabetes, based on family history.
- **Age**  
Age of the patient in years.
- **Outcome**  
Binary target variable:
  - 1 = patient is diagnosed with diabetes
  - 0 = patient is not diagnosed with diabetes

Your task is to build and analyse binary classification models that predict **Outcome** from these clinical features. You will first clean and prepare the data in KNIME, then train a

decision tree model, evaluate its performance, and explore global feature importance to understand which variables contribute most to the diabetes prediction.

## KNIME Instructions



### 1) Data Preparation block

**1.1 CSV Reader** – Drag CSV Reader, connect nothing to it, double-click, and set the path to **diabetes.csv** so KNIME can read the Pima Diabetes data.

**1.2 Statistics (raw)** – Drag Statistics, connect it to the CSV Reader output; execute and open the view to quickly inspect distributions and spot weird values (0's, outliers, class balance).

**1.3 Number to String (Outcome)** – Drag Number to String, connect from CSV Reader, configure it so **Outcome** is converted from integer to string; this gives KNIME a proper *nominal* class column for classification nodes.

#### 1.4 Row Filter (drop-0s demo)

Drag a **Row Filter** node, connect it after **CSV Reader**, and set criteria such as BMI > 0, Glucose > 0, BloodPressure > 0, SkinThickness > 0, and Insulin > 0 with “Match row if matched by: All criteria” so KNIME keeps only records with non-zero medical values, demonstrating a naïve “drop rows with 0s” cleaning approach.

**Row Filter** [X]

Click to close

**Filter**

Match row if matched by

☒ All criteria ☐ Any criterion

**Criterion 1** [Up] [Down] [Delete]

Filter column: BMI Operator: Greater than

Value: 0

**Criterion 2** [Up] [Down] [Delete]

Filter column: Glucose Operator: Greater than

Value: 0

### 1.5 Statistics (show data loss from drop-0s)

Drag a **Statistics** node, connect it after this Row Filter, execute, and open the view to see that the row count falls from 768 to only ~400 rows, illustrating that this hard filtering discards many patients and motivating the other branch, where we convert 0s to missing values and impute them instead.

### 1.6 Expression (force 0 values to missing)

Drag an **Expression** node, connect it after **Number to String**, and for each clinical column (Glucose, BloodPressure, SkinThickness, Insulin, BMI) add an expression like `if($Glucose$ == 0, MISSING, $Glucose$)` (set “Output replaces ‘Glucose’”) so that any impossible 0 is turned into a missing value that can later be filled by the **Missing Value** (mean imputation) node instead of dropping whole rows.

Expression 1

↑

↓

📄

🗑️

1

if(\$['BloodPressure'] == 0, MISSING, \$['B

▽ Output replaces "BloodPressure"

Append

Replace

📄 BloodPressure

▽

Expression 2

↑

↓

📄

🗑️

1

if(\$['BMI'] == 0, MISSING, \$['BMI'])

> Output replaces "BMI"

### 1.7 Missing Value (mean imputation for numeric columns)

Drag a **Missing Value** node, connect it after **Expression**, then double-click to open its configuration and on the **Default** tab set Number (Integer), Number (Long Integer) and Number (Float) to **Mean** (leave String as “Most Frequent Value”); this fills all the new missing values in the medical numeric columns with their column means so the next nodes receive a complete table with no missing entries.

Number (Integer)	Mean
Number (Long Integer)	Mean
Number (Float)	Mean
String	Most Frequent Value

**1.8 Table Partitioner** – Drag Table Partitioner, connect from **Missing Value**, choose a train/test split (e.g. 70/30) and tick “Stratified” by **Outcome**; this creates a fair split with similar class balance in both partitions. To make your results reproducible, enable **Fixed random seed** and set its value to **2026**.

Table Partitioner

×

First partition type

Relative (%)

Absolute

Relative size

70

^

v

Sampling strategy

Random

Stratified

Linear

First rows

☒ Fixed random seed

2026

^

v

If input table is empty

Fail

Output empty table(s)

## 2) Decision Tree Model block

2.1 **Decision Tree Learner** – Drag Decision Tree Learner, connect its data input to the output port of Table Partitioner (training data), configure **Outcome** as class column, and other settings (Gini, pruning, etc.), and execute to train the baseline tree.

Dialog - 4:9 - Decision Tree Learner (Train Decision Tree)

Options PMMLSettings Flow Variables Job Manager Selection

General

Class column S Outcome

Quality measure Gini index

Pruning method MDL

☒ Reduced Error Pruning

Min number records per node 10

Number records to store for view 10,000

☒ Average split point

Number threads 8

☒ Skip nominal columns without domain information

Root split

☐ Force root split column

Root split column D Age

Binary nominal splits

☒ Binary nominal splits

Max #nominal 10

☐ Filter invalid attribute values in child nodes

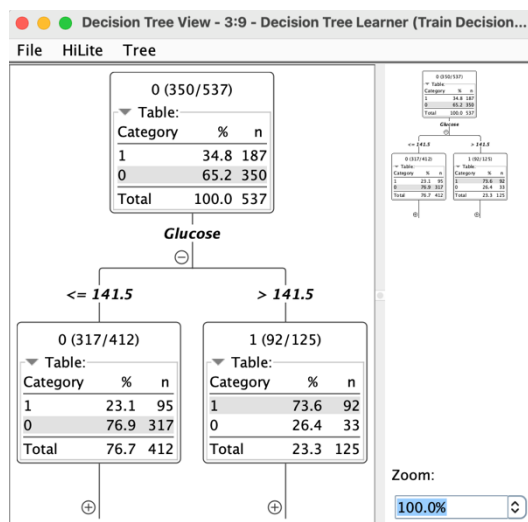
OK

Apply

Cancel

?

You can inspect the learned Decision Tree by **right-clicking the Decision Tree Learner node** → **Open view**. This opens a window showing the full tree structure, including each split (feature and threshold) and the class distribution at every node



**2.2 Decision Tree Predictor (direct KNIME evaluation)** – Drag Decision Tree Predictor, connect its model input to the Decision Tree Learner and its data input to the output port of Table Partitioner (test data); execute to get predictions.

**2.3 Scorer** – Drag Scorer, connect from Decision Tree Predictor output and configure Outcome as true class; execute to view accuracy, confusion matrix, ROC etc. – this is the KNIME-side evaluation of the baseline.

#	RowID	TruePositiv... Number (Int...)	FalsePositi... Number (Int...)	TrueNegati... Number (Int...)	FalseNegat... Number (Int...)	Recall Number (Flo...)	Precision Number (Flo...)	Sensitivity Number (Flo...)	Specificity Number (Flo...)	F-measure Number (Flo...)	Accuracy Number (Flo...)
1	1	48	29	121	33	0.593	0.623	0.593	0.807	0.608	⚠
2	0	121	33	48	29	0.807	0.786	0.807	0.593	0.796	⚠
3	Overall	⚠	⚠	⚠	⚠	⚠	⚠	⚠	⚠	⚠	0.732

**2.4 PMML Writer (export tree to Python)** – Drag PMML Writer, connect the model port from Decision Tree Learner, and configure it to write d-tree.pmml into the destination folder; this file will later be loaded and evaluated in Python.

## 2.5 Column Filter (simulate unlabeled deployment data)

Drag a **Column Filter** node, connect it from the Table Partitioner (**test dataset**), and *deselect* the **Outcome** column so only the predictor features remain; this simulates a real

deployment situation where you want to score new patients but do not know their diagnosis yet.

## 2.6 PMML Reader + Decision Tree Predictor (deployment branch)

Use **PMML Reader** to load the saved **d-tree.pmml**, then drag a second **Decision Tree Predictor** and connect its *model* input to the PMML Reader and its *data* input to the Column Filter output; this shows how a trained tree model, stored as PMML, can be reloaded and applied to feature-only data in a separate workflow or at deployment time.

## 2.7 CSV Writer (prediction.csv for external use)

Drag a **CSV Writer**, connect it to the second Decision Tree Predictor, and configure it to write prediction.csv into the destination folder; this exports the predicted class and probabilities for each patient, which can be used as the final deployment output.

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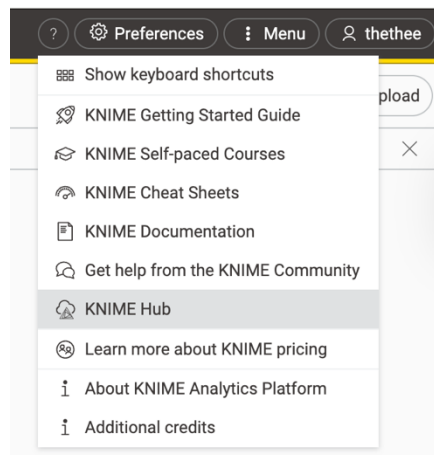
## 3) Decision Tree Model & Global Feature Importances block

**3.1 Capture Workflow Start / End (wrap LR workflow as object)** – Drag Capture Workflow Start before the **Decision Tree Predictor** and Capture Workflow End after them; this turns the whole **Decision Tree** scoring workflow into a **workflow object** that the Global Feature Importance component can call internally.

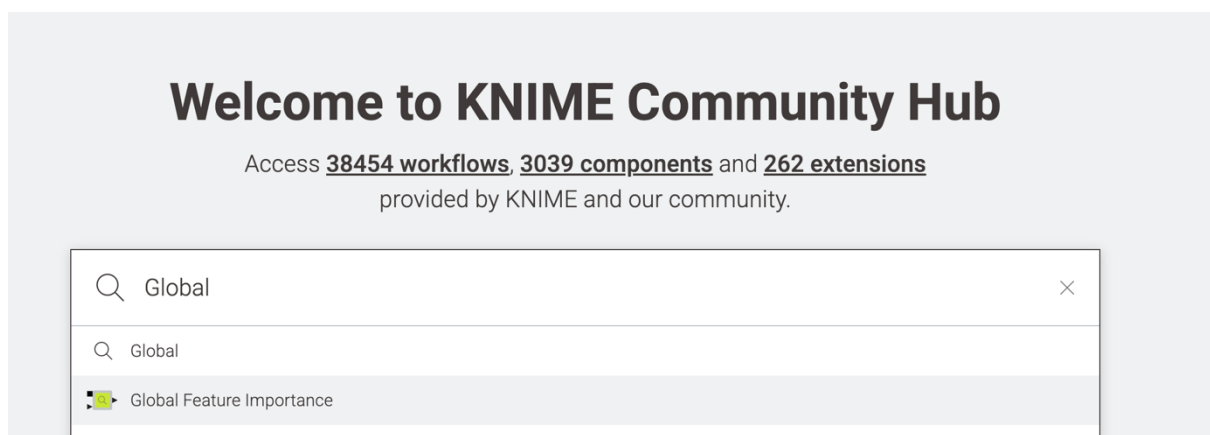
**3.2 Decision Tree (2) Predictor** – Drag **Decision Tree (2) Predictor**, connect the model input from the **PMML Reader** and the data input from the output of **Capture Workflow Start**, then connect its output to **Capture Workflow End**.

**3.3 Global Feature Importance (download component from KNIME Hub and connect it)**

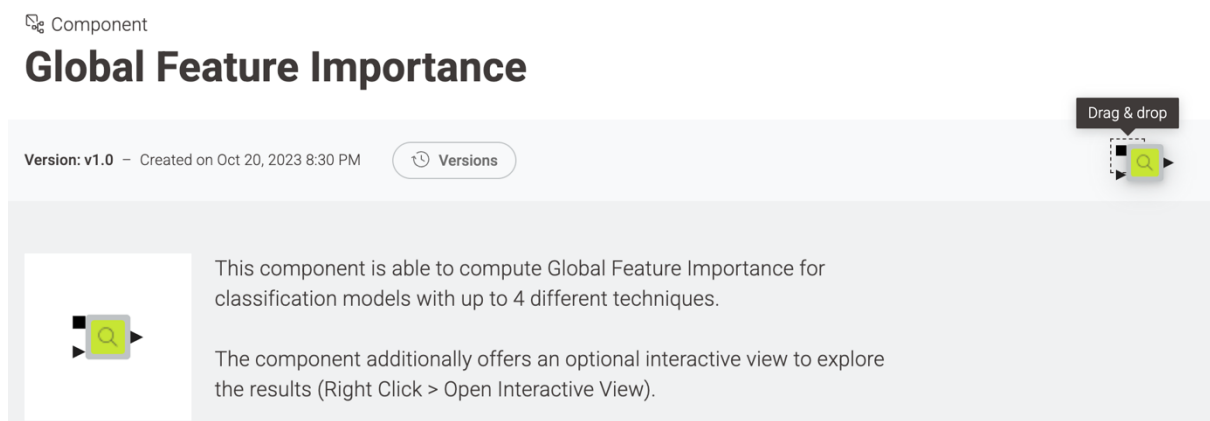
At the top-right of KNIME, click '(?) button' → KNIME Hub,



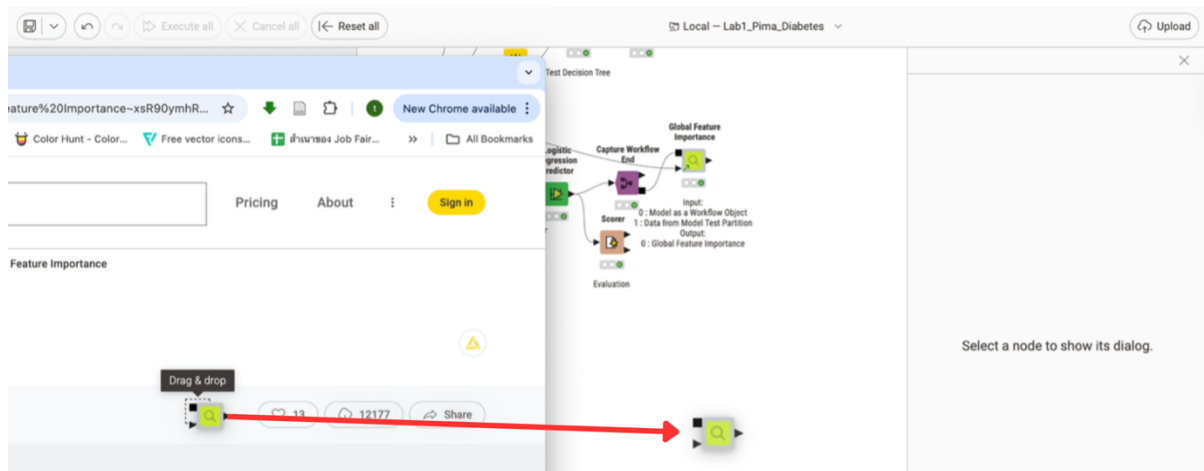
then in the Hub search box type “**Global Feature Importance**” and open the component page;



on that page, grab the small green **Drag & drop** icon and drag it straight into your workflow editor,







then connect its **model workflow** input port 0: Model as a Workflow Object to **Capture Workflow End**, its **data** input port 1: Data from Model Test Partition to the **second output of Table Partitioner** (test data), execute the node, and finally right-click → **Open Interactive View** to explore how each feature contributes to the **Decision Tree** model.

Global Feature Importance

Target column and focus class:

Column

Outcome

Value

1

Importance methods:

☒ Surrogate Generalized Linear Model

☒ Surrogate Decision Tree

☒ Surrogate Random Forest

☒ Permutation Feature Importance

Performance metric:

F-measure

For the method "Permutation Feature Importance..."

1

Show top n features:

10

Surrogate models data pre-processing: maximum...

100

Discard

Apply and Execute

Apply

Below is an example of the Global Feature Importance output. To view it in KNIME, right-click the **Global Feature Importance** node and choose **Open View**.

