# Interpretation Tree Model Matching

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# Interpretation Tree Model Matching

#### Given:

Sets of model lines  $\{m_i\}$  in a scene coordinate system Set of image lines  $\{d_i\}$  in an image coordinate system

Image to scene scale conversion factor  $\sigma$  (pixels to cm)

#### Do:

- 1. Match image and model lines  $\{(m_i, d_j)\}$
- 2. Estimate transformation mapping model onto data: R,  $\vec{t}$
- 3. Verify matching and pose estimate

Output: identity and position  $(R, \vec{t})$ 

## Interpretation Tree matching

Goal: Correspondence between subset of M model features  $\{m_i\}$  and D data features  $\{d_j\}$ 

Complete (exhaustive, depth-first) search - if a match exists, it will be found

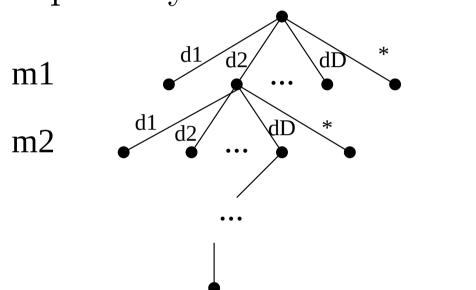
Needs a 'wildcard' ('\*') data feature to match model features with no corresponding data feature (occlusion, segmentation failure)

Can find multiple solutions

**Result**:  $\{(m_i, d_{j_i})\}$  set of matched features

#### Search Tree

Expand by model feature at each new level



mM d1 d2

Any given node in tree represents a set of matches  $\{(m_i, d_{j_i})\}$ 

# Reducing Search Complexity

Do we need to consider all paths in search tree? No: Suppose current match state has these pairs matched:  $\{(m_i, d_{j_i})\}, i = 1..k$  Given a new pair  $(m_{k+1}, d_{j_{k+1}})$ 

- 1.  $unary\_test(m_{k+1}, d_{j_{k+1}})$  terminates extending search path if new pair has incompatible properties
- 2.  $binary_test(m_{k+1}, d_{j_{k+1}}, m_x, d_{j_x})$  for all x = 1..k terminates extending search path

if new pair has incompatible properties with each previous pairing on this tree branch (as all parts of the same object are compatible).

- 3. Early success limit L can stop search when have  $\{(m_i, d_{j_i})\}, i = 1..L$  compatible pairs
- 4. Early failure limit L can stop search when can never get L pairs on this path. If have t non-wildcard matches on this path out of k pairings, then fail if t + (M k) < L

## Computational Complexity

M model feature tree levels. D data features on each level plus 1 wildcard

Worst case:  $(D+1)^M$  nodes in tree to visit

 $p_u$  - probability that any random model feature and any random data feature pass unary\_test

 $p_b$  - probability that any 2 random model features and any 2 random data features pass binary\_test

Then, if  $p_b MD < 2$ , then the average case complexity of ITREE search is  $O(LD^2)$ 

Much smaller, but can still be substantial

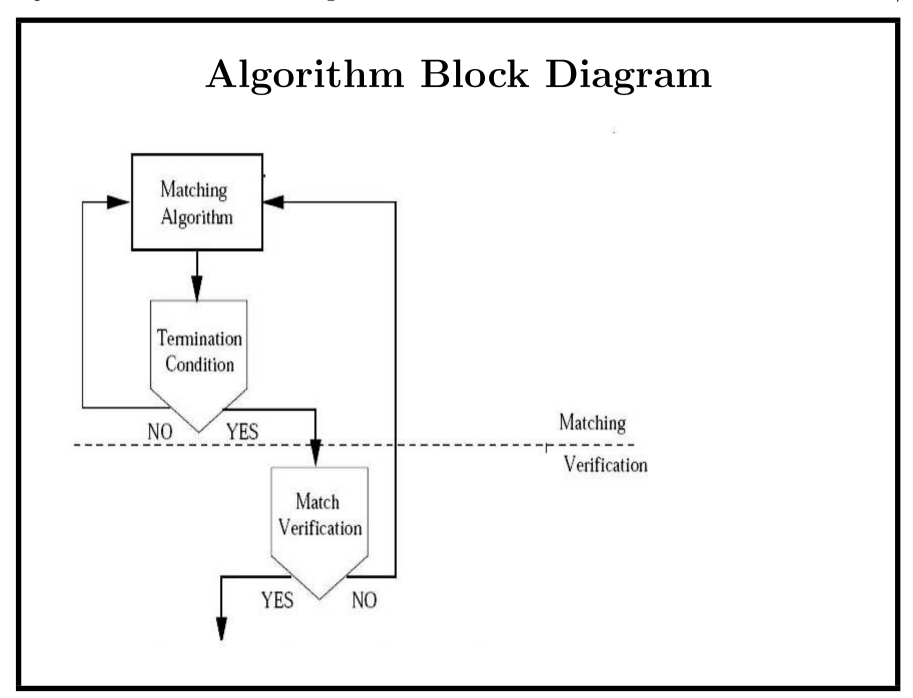
#### IT algorithm matlab code

```
% interpretation tree - match model and data lines until
% Limit are successfully paired or can never get Limit
% model - current model
% numM - number of lines in the model
% mlevel - last matched model feature
% Limit - early termination threshold
% pairs(:,2) - paired model-data features
% numpairs - number of paired features
function ok=itree(model,numM,mlevel,Limit,pairs,numpairs)
global Models numlines datalines
% check for termination conditions
```

```
[theta,trans] = estimatepose(model,numpairs,pairs)
  for p = 1 : 4
    ok = verifymatch(theta(p),trans(p,:)',model,
              numpairs,pairs);
    if ok
                % successful verification
      return
    end
  end
                % failure to verify - continue search
  return
end
% never enough pairs
if numpairs + numM - mlevel < Limit
  ok=0;
  return
end
```

```
% normal case - see if we can extend pair list
mlevel = mlevel+1;
% do unary test
 if unarytest(model,mlevel,d)
   % do all binary tests
   passed=1;
   for p = 1: numpairs
     if ~binarytest(model,mlevel,d,pairs(p,1),pairs(p,2))
       passed=0;
       break
    end
   end
```

```
if passed
      % passed all tests: add to matched pairs and recurse
      pairs(numpairs+1,1)=mlevel;
      pairs(numpairs+1,2)=d;
      ok=itree(model,numM,mlevel,Limit,pairs,numpairs+1);
      if ok
               % successful verification
        return
      end
    end
  end
end
% wildcard case - go to next model feature
ok = itree(model,numM,mlevel,Limit,pairs,numpairs);
```



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# Line matching unary test

DATA LINE MODEL LINE

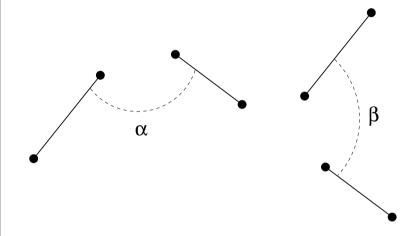


Pass test if  $\sigma l_m(1 - \delta_u) \leq l_d \leq \sigma l_m(1 + \delta_u)$ 

Allows for calibration and segmentation errors Position independent property ( $\delta_u = 0.3$  typical)

## Line matching binary tests

DATA LINES MODEL LINES



Pass test if  $|\alpha - \beta| \leq \delta_b$ 

Allows for calibration and segmentation errors Position independent property ( $\delta_b = 0.2$  radians typical)

Also: don't allow duplicate use of model or data lines

#### Matching performance

Limit L = number of model lines - 1 Tries all models Stops at first verified model instance for each model

#### Different Matched Models & Instances

Image	True Model	Tee	Thin L	Thick L
1	Tee	4	0	12
2	Tee	4	0	12
3	Tee	21	0	12
4	Tee	21	0	12
5	Thin L	0	15	2
6	Thin L	0	15	2
7	Thin L	0	15	2
8	Thin L	0	24	2
9	Thick L	0	2	3
10	Thick L	0	2	3
11	Thick L	0	2	3
12	Thick L	0	2	3

#### What Have We Learned?

Introduction to

- General Feature Matching Algorithm
- Efficient if good unary/binary tests
- Suitable for 50% (estimated) flat parts