Comparison between a sequential and a multithreading version of the mean shift clustering algorithm

Emilio Cecchini

emilio.cecchini@stud.unifi.it

Abstract

In this paper, after a brief introduction to the mean shift clustering, a sequential version will be compared with a multithreading version of the algorithm. The obtained speedup with a multi-core CPU will be analyzed using a different number of threads. The algorithm is written in C++ and the parallel version is obtained with OpenMP. The focus of this paper and his associated code do not consist in showing a very efficient version of the mean shift clustering algorithm, but rather in analyzing the performance improvements obtainable with a multithreading version compared to a sequential one.

1. Introduction

Since there are many variations of the mean shift algorithm, in this section I will describe the specific version used in my implementation.

1.1. Mean shift

The mean shift algorithm is a nonparametric clustering technique that does not require as input the number of clusters to look for. It is based on the concept of *kernel density estimation* or *KDE*, that is a method to estimate the underlying distribution for a set of data.

At each step, a *kernel function* is applied to each point that causes the points to shift in the direction of the local maxima determined by the kernel. The iterations end when all points reach the maxima of the underlying distribution estimated by the chosen kernel.

There are many different types of kernel, the most used is the *Gaussian kernel*:

$$k(x) = e^{-\frac{x}{2\sigma^2}} \tag{1}$$

The standard deviation σ is the *bandwidth* parameter. Depending on the kernel bandwidth parameter used, the

resultant density function will vary: with a high bandwith value you will get a few large clusters and vice versa.

The new location where to shift each point at each step of the algorithm is computed as a weighted average between the point and its neighbors, where the weights are calculated with the Gaussian kernel. Suppose x is a point to be shifted and N(x) are the sets of points near to that point. Let $dist(x,x_i)$ be the distance from the point x to the point x_i . The new position x' where x has to be shifted is computed as follows:

$$x' = \frac{\sum_{x_i \in N(x)} k(dist(x, x_i)^2) x_i}{\sum_{x_i \in N(x)} k(dist(x, x_i)^2)}$$
(2)

The mean shift algorithm applies that formula to each point iteratively until they converge, that is until the position does not change.

1.2. The algorithm

The algorithm is very simple: each point shifts towards the maxima of its underlying distribution. The algorithm ends when all the points have stopped shifting.

Here is the pseudocode of the core of the algorithm:

```
Algorithm 1 Mean shift core

while allPointsHaveStoppedShifting() do

for each point p do

if hasStoppedShifting(p) then

continue

end if

shift(p)

end for
end while
```

To speed up the process, the shifting process of a point is stopped when the distance from its older position is less than an epsilon value specified by the user.

1.3. The parallel version

The mean shift algorithm is a embarrassingly parallel work: each point perform its shifting independently from

the other points. This makes it the perfect case for using the OpenMP technology. In fact, with a single pragma command it was possible to switch from a sequential version to a parallel version.

Here is the pseudocode of the parallel version of the core algorithm:

Algorithm 2 Mean shift core parallel

```
while allPointsHaveStoppedShifting() do
    #pragma omp parallel for schedule(dynamic)
    for each point p do
        if hasStoppedShifting(p) then
            continue
    end if
        shift(p)
    end for
end while
```

Note that the only difference from the sequential version in 1.2 is the pragma statement. That statement is placed just before the for loop, in this way there is no need of any critical sections. If it had been placed before the while loop, then the parallel algorithm would have been more complex introducing an overhead due to the synchronization between threads.

The clause schedule (dynamic) is important because at the end of the computation, there will be some points that have stopped shifting, so they immediatly end the computation of the for loop. With the dynamic scheduling there will be a better distribution of the workload of each thread.

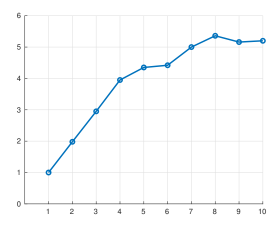
1.4. Speedup analysis

The speedup is computed with a script that first executes a sequential version, then it executes on the same data set differents parallel versions using a different number of threads.

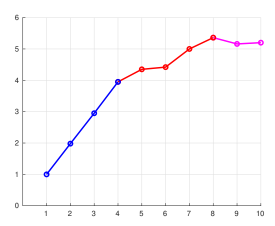
Here is a table and a chart reporting the speedup of the parallel version with a different number of threads. The test was performed on a machine with an Intel Xeon CPU 2.00 *GHz* with 8 cores on a data set of 10000 points of three dimensions:

Sequential version: 54.1719 s

Threads	Time (seconds)	Speedup
2	27.42	1.98
3	18.36	2.95
4	13.72	3.95
5	12.44	4.35
6	12.25	4.42
7	10.83	5.00
8	10.11	5.36
9	10.50	5.16
10	10.42	5.20



The speedup seems to grow with three different patterns. From two threads to four threads the speeup is almost perfectly linear, from five to eight it seems to be sub-linear and over eight threads it stops growing.



The explanation for these three different growing trends is immediate. The machine where the test was performed has the hyperthreading technology enabled: it shows eight cores but actually it has only four physical cores, the other four cores are virtual. In fact, from two to four threads the

speedup is linear because each physical core has to execute a single thread. Using from five to eight threads can speed up the execution, but the speedup is no more perfectly linear, because there are more threads than physical cores. Finally over eight threads there is no more improvements in terms of speeds, on the contrary, it is slightly slower due to the overhead of the contex switches between threads.