# Theoretical Cost Comparison of Remote Visualization Strategies

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#### Abstract

We compare three remote visualization strategies used for interactive exploration of large data sets in distributed environments: image-based rendering, parallel visualization servers, and subsampling. To determine the regimes for which each approach is most cost-effective, we develop performance models to study the computation and communication costs associated with the common visualization task of isosurface generation. For one particular strategy, subsampling, we further investigate the tradeoffs between multiresolution and uniform grid methods in terms of performance and approximation errors.

Keywords: Remote Visualization, Multiresolution Subsampling, Performance Models.

### **1** Introduction

Interactively exploring tera- and petabyte data sets is extremely challenging, particularly for scientists whose primary access to visualization resources is a desktop graphics workstation. To address this problem, researchers are exploring approaches that combine remote, high-end resources with high-speed networks to provide interactive navigation and exploration of very large data sets.

In this paper, we consider three common strategies for improving performance of interactive data exploration using remote resources: image-based rendering, parallel visualization servers, and subsampling of the original data set. We give an overview of these strategies in Section 2. Each approach attempts to improve performance by targeting a different stage of the visualization process, and each has its advantages and disadvantages. For this analysis we consider each method individually, but note that they are not inherently exclusive and combinations are possible. To determine the most cost-effective strategy for particular hardware system or problem configurations, we examine the performance characteristics of each by developing theoretical performance models. These models estimate the costs of computation and communication when parameters such as network bandwidth, problem size, and visualization demands change. We describe the models in Section 3 and analyze them in Section 4 for a variety of problem scenarios.

For the subsampling approaches, we develop two theoretical models: one for uniform grids and one for multiresolution grids. Uniform grids are advantageous for two reasons: (1) they can be represented with a comparatively small amount of data, and (2) visualization algorithms on uniform grids generally outperform their nonuniform grid counterparts. Therefore, a larger number of grid points can be used with uniform grid subsampling than with multiresolution techniques. However, multiresolution techniques are designed to minimize the approximation error associated with subsampling by placing more points where the data is changing rapidly. These tradeoffs are examined in more detail for two different application data sets in Section 4.

## 2 Remote Data Exploration Strategies

For each of the three strategies considered for remote data exploration, we briefly describe the fundamental concepts and give an overview of the method's advantages and disadvantages.

**Image-based rendering** techniques use two or more reference images from multiple viewpoints to reconstruct either the geometry in a scene or new images of the scene as the user's viewpoint changes. In this paper, we focus on image warping techniques that use reference images containing color, depth, and surface normal information to "warp" or change the input images to the desired output image (e.g., [1, 13]). Typically, several reference images are needed to avoid approximation errors or holes when the scene is reconstructed from an arbitrary viewpoint. The primary advantage of this technique for remote data exploration is that the amount of data transmitted and manipulated locally is independent of the complexity of the scene or original data set. Thus the communication and local computation costs are fixed as data set sizes increase. In addition, if remote resources are used to generate the derived visualization entities using the full data set, no approximation errors are associated with the reference images. However, as the user changes the view perspective, errors associated with the reconstruction process can misrepresent the original data set. In addition, these techniques are still moderately expensive, and even fairly sophisticated hierarchical techniques can require 1.5 seconds/frame using four reference images [1].

**Parallel visualization servers** utilize remote computational resources to visualize full-resolution data sets either as the computation proceeds (e.g., [5, 16]) or as a post-processing step (e.g., [2, 6]). The geometries of the derived visualization entities, rather than images, are extracted and communicated to the graphics workstation for display. Typically, lower-dimensional entities such as isosurfaces or streamlines are targeted for use with these systems because their transmission and memory requirements are much smaller than the full-dimensional data set. The primary advantage of these techniques is that the derived visualization entities have no subsampling or other approximation errors. In addition, once the geometry is loaded into the local workstation, it may be freely manipulated without reconstruction errors. However, the size of the geometries transmitted to the local graphics workstation are functions of the overall problem size. Thus, as the problem size increases, the demands on computational resources, both remote and local, and on network bandwidths also increase.

**Subsampling and clustering** techniques create smaller, full-dimensional data sets by sampling the original data at specified locations or by averaging clusters of points from the original data set. The simplest approach to subsampling is to create a uniform grid representation of the original data set. Alternatively, a hierarchical, multiresolution representation of the data can be constructed using, for example, quadtrees or octrees [10, 9], progressive meshes [8], wavelets [14], or other clustering approaches [7]. The level of detail in each region is controlled through a variety of mechanisms, such as error tolerance bounds that control fidelity to the original model, or user input, such as field of view. Subsampling or clustering approaches are useful for fast, local exploration of the reduced data set. The remote computation and transmission costs are fixed regardless of the number of visualization tasks. The primary disadvantage is the difficulty of maintaining fidelity

to the original data set. As the original problem size increases, a smaller percentage of points can be used in the reduced data set resulting in higher approximation errors. This disadvantage can be somewhat mitigated, at the cost of more communication, by an adaptive windowing approach. Furthermore, forming derived visualization entities on the interpolated data set may be difficult in the multiresolution approach; for example, isosurfaces may contain undesired "cracks."

### **3** Performance Models

For each remote data exploration strategy, we now develop a theoretical performance model describing the computation and communication costs. The visualization task used in these models is isosurface generation, which we chose for three reasons: (1) Isosurfaces are one of the most commonly used visualization techniques for exploring scientific data sets. (2) Much research in the visualization community has targeted efficient isosurface generation for both uniform and multiresolution data sets and for both serial and parallel computers. (3) The fact that isosurfaces are lower-dimensional entities ensures that all three strategies are fairly considered. Other commonly used visualization techniques such as cutting planes, streamlines, vector glyphs, and volume visualization could be easily modeled by replacing the isosurface-specific information. We note that the performance of the parallel visualization server strategy can be dramatically affected by the choice of visualization techniques studied. For example, the cost of transmitting streamline information is very small and would likely improve the relative performance of the parallel visualization server compared to the other two strategies.

The costs in our models include the time to compute a specified number of isosurfaces, the time to transmit information over a wide area network to the local graphics workstation, and, for the subsampling techniques, the time to compute the reduced data sets. Because the data sets of interest are large, we assume that the scientist has access to a remote parallel computer and that any remote isosurface or subsampling computations are done scalably in parallel. We assume that the original data is preloaded and distributed across the processors of the remote parallel computer, because this is common to each approach and does not differentiate the models. For all models, we assume that the original computational mesh is nonuniform.

We note that our models serve to highlight the basic performance of each remote visualization strategy and some simplifying assumptions have been made. First, we do not address the costs associated with loading and processing the resulting data sets on the local graphics workstation. Second, in the case of parallel visualization servers, the number of primitives contained in the isosurface may be reduced through various decimation procedures (see, for example, [3]); we do not include these strategies in our models. Finally, our models do not address the general question of compression of the communication between the remote and local computers.

The parameters and cost variables used in our models are defined in Table 1. To define the hardware characteristics of our remote visualization system, we use P to define the number of remote processors available for parallel computation and R and L to define the network bandwidth and latency in Mbs and seconds, respectively. The number of elements in the data set, N, and the number of isosurfaces to be computed and rendered, I, define the complexity of our visualization task. The parameter X defines the image size in pixels for image-based rendering, and  $N_R$  defines the number of elements in the reduced data sets for subsampling. The parameters  $C_{R_u}$  and  $C_{R_m}$  give the total serial cost of subsampling for uniform and multiresolution grids, respectively. The parameters  $C_{I_u}$  and  $C_{I_m}$  give the cost per element of isosurface generation on uniform and

multiresolution grids, respectively. A detailed analysis of these costs will be given in Section 4.

General Costs		System-Dependent Computational Costs				
P	Number of Remote Processors	$C_{R_u}$	Uniform Grid Subsampling Cost			
R	Network Bandwidth (Mbs)	$C_{R_m}$	Multiresolution Subsampling Cost			
L	Network Latency (s)	$C_{I_u}$	Uniform Grid Isosurface Cost/Element			
N	Number of Elements	$C_{I_m}$	Multiresolution Isosurface Cost/Element			
Ι	Number of Isosurfaces					
X	Pixels/Image					
$N_R$	Number of Subsampled Elements					

Table 1: Cost model variables

#### 3.1 Model 1: Image-based Rendering

For image-based rendering techniques, we assume that the isosurfaces are computed in parallel and that six depth images are used for reconstruction on the local graphics workstation [13]. To determine the number of bits that must be transmitted for each pixel in the depth image, we use the best-case scenario information given in [11] for a postrendering warping technique. In particular, we assume 24 bits for color, 16 bits for depth, and 8 bits for surface orientation information, for a total of 48 bits per pixel. We assume that a new set of reference depth images is required for each new isosurface generated. The total cost of the image-based rendering technique is

$$M_I = \frac{N C_{I_m} I}{P} + \frac{48 \cdot 6 X I}{10^6 R} + 2 L I.$$
(1)

The first term gives the time required to compute the isosurfaces in parallel on the original data set. The second term gives the transmission time required to send the depth images associated with each isosurface. The final term gives the network latency for new isosurface requests. We assume that the cost of generating the six depth images is negligible and the amount of information needed to request new isosurfaces is minimal.

#### 3.2 Model 2: Parallel Visualization Servers

The parallel visualization server also computes the isosurfaces in parallel, so the first term of this model is identical to that in Equation 1. The data transmitted for each isosurface are three spatial coordinates for each data point (3 32-bit floats) as well as the connectivity information for each triangle (3 32-bit integers). Using the fact that the numbers of vertices and elements in a triangular mesh are nearly equal and that the average isosurface will contain  $N^{\frac{2}{3}}$  triangles, the total cost of the parallel visualization server is

$$M_V = \frac{N C_{I_m} I}{P} + \frac{32 \cdot 6 N^{\frac{2}{3}} I}{10^6 R} + 2 L I.$$
<sup>(2)</sup>

#### 3.3 Model 3: Uniform Subsampling

For uniform grid subsampling, we assume that the original data is partitioned such that the reduction operations may be performed with minimal communication and will scale linearly as a function of P. The transmission costs for a uniform grid include the cost of sending the origin, grid spacing, and problem size in each of the three dimensions (9 floats), and the scalar data and error associated with each element  $(2 \cdot N_R \text{ floats})$ . No additional spatial or coordinate information is required. The total cost of uniform subsampling is

$$M_U = \frac{C_{R_u}}{P} + N_R C_{I_u} I + \frac{32 \cdot (9 + 2N_R)}{10^6 R} + 2L.$$
(3)

The first two terms are the computational costs associated with data reduction and isosurface generation on the reduced uniform grid, respectively. The third and fourth terms are the transmission and latency costs for a single request for a subsampled grid, respectively.

#### 3.4 Model 4: Multiresolution Subsampling

There are many approaches for multiresolution subsampling, and we develop our cost model by examining one particular approach that uses parallel octrees. In this method, points from the original data set are inserted into the appropriate leaf octants. These are then evaluated according to a specified criterion and refined if necessary with associated data points reassigned to the new leaf octants. When all data has been inserted, an average data value is computed for each leaf, and this constitutes the reduced data set. The user may interactively change the leaf criterion, either globally or in a specified region of interest, and thereby the resolution of the reduced data. The criterion may also be generated automatically to meet a performance constraint such as maximum target number of leaves. To provide an indication of the error associated with the reduced data set, we compute normalized standard deviation,  $\sigma_n$ , and maximum deviation from the mean,  $e_n$ , for each leaf octant. These values also serve to highlight potential regions of interest; large deviations from the average value may indicate fine-scale structure that was not adequately captured by the reduction process. Additional details about the parallel octree algorithms, our software architecture, and results obtained on large data sets can be found in [4].

The model for multiresolution subsampling is similar to Equation 3. Because we are using an octree data representation, the spatial coordinates and connectivity information for the reduced mesh must be transmitted in addition to the scalar information. Given a number of octant leaves, the number of associated vertices cannot be determined *a priori*. An upper bound for the number of vertices is  $8N_R$ , which is the case when no vertex is shared between octants. A lower bound for the number of vertices is  $N_R$ , which is the case when the vertices are maximally shared by octants, that is, when the leaf octants form a uniform mesh. For the purposes of our model, we assume an average case of  $4N_R$ . Thus, the total amount of information that must be transmitted is  $3 \cdot 4N_R$ floats for the spatial coordinates plus  $8N_R$  integers for the connectivity data and  $2N_R$  floats for the scalar and error data. With these assumptions, our cost model for multiresolution subsampling is

$$M_M = \frac{C_{R_m}}{P} + N_R C_{I_m} I + \frac{32 \cdot (22N_R)}{10^6 R} + 2L.$$
(4)

The primary difference between Models 3 and 4 is the amount of data that must be transferred for the reduced data set, and the use of the nonuniform grid costs for data reduction and isosurface generation,  $C_{R_m}$  and  $C_{I_m}$ , in the first two terms of the model.

### **4** Results

To determine the regimes for which each of the models developed in Section 3 are most costeffective, we first determine typical values for the cost parameters  $C_{I_u}$ ,  $C_{I_m}$ ,  $C_{R_u}$ , and  $C_{R_m}$ . We then analyze the performance models for various values of N, P, I, and R. Finally, we consider the uniform grid and multiresolution subsampling techniques in more detail and examine their performance as a function of the subsampling error for two different application problems. All experiments were performed on one or more processors of an SGI Origin with 250 MHz R10000 chips, and all timings were performed using the Unix subroutine gettimeofday().

Network bandwidth and latency estimations are based on the vBNS and ESnet networks. Each of these networks consists of OC3 lines for a maximum throughput of 155 Mbs, although expected performance is often far less. Latency measurements listed on the vBNS net traffic web page [12] range from 3 to 40 ms depending on the destination/origination. For the purposes of this paper, we consider bandwidth ranging from R=.5 to 100 Mbs and use a latency of L = 20 ms.

#### 4.1 Determining Cost Parameter Values

We first determine the cost per element of isosurface generation on uniform and multiresolution grids,  $C_{I_u}$  and  $C_{I_m}$ , respectively. We use vtk's vtkContourMarchingFilter, which uses a fast marching cubes algorithm for uniform structured point sets and a general algorithm for all other mesh types, including our unstructured octree representation [15]. We tested the algorithms on the same uniform data set, changing only the way it was represented in vtk data structures. In particular, we used vtkStructuredPoints to determine  $C_{I_u}$  and vtkUnstructuredGrid to determine  $C_{I_m}$ . Timings were conducted for five different problem sizes ranging from  $N = 20^3$  to  $60^3$  to determine the cost per element for computing the isosurface. As N increases,  $C_{I_u}$  asymptotically approaches a value of approximately .0015 ms/cell, and this is the value used in our model. In contrast, the cost of using general data structures in vtk, along with the associated virtual function calls, results in the parameter  $C_{I_m}$  remaining fixed at approximately .024 ms/cell, a factor of 16 greater than  $C_{I_u}$ . We note this factor would be less for routines specific to octree data types and is also implementation-dependent.

To determine the uniform subsampling cost,  $C_{R_u}$ , we subsampled a tetrahedral mesh onto a uniform grid of different sizes and monitored the time required to insert the data and compute the average value and normalized standard deviation for each subsampled grid point. We also include the time required to compute the average and maximum deviation over the entire reduced data set. Timings were conducted for  $N_R = 1$ ,  $5^3$ ,  $10^3$ , and  $20^3$ . Linear least-squares analysis for  $N_R = 1$ yields a base cost for visiting every point in the original data set of  $8.75 \cdot 10^{-7}N$  seconds. The total data reduction costs grow as a function of both N and  $N_R$ , and subsequent least-squares analyses yield the final expression for  $C_{R_u}$ :

$$C_{R_u} = 8.75 \cdot 10^{-7} N + (6.50 \cdot 10^{-5} + 4.31 \cdot 10^{-12} N) N_R.$$

To determine the multiresolution subsampling cost,  $C_{R_M}$ , we performed a similar analysis using the octree data reduction technique described in Section 3.4. Linear least-squares analysis for the  $N_R = 1$  case yields a base cost of  $3.21 \cdot 10^{-5}N$  seconds. Again, data reduction costs increase as a function of N and  $N_R$ . Linear least-squares analyses yield our final formula for  $C_{R_m}$ :

$$C_{R_m} = 3.21 \cdot 10^{-5} N + (.87 \cdot 10^{-5} + 2.5 \cdot 10^{-10} N) N_R$$

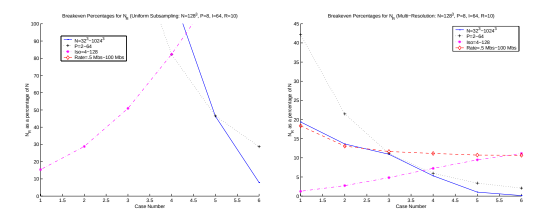


Figure 1: The breakeven graphs for uniform and multiresolution subsampling with base parameters  $N = 128^3$ , P = 8, R = 10 Mbs, I = 64.

#### 4.2 Performance Model Comparisons

To determine the regimes for which each model is the most cost-effective, we first consider the image-based rendering and the parallel visualization server models, Models 1 and 2, respectively. The computational costs for isosurface generation are identical; the models are differentiated only by the amount of data transmitted. By equating the two models, we derive the breakeven function that relates the number of pixels in the image, X, and the data set size, N:

$$X = \frac{2}{3}N^{\frac{2}{3}}.$$
 (5)

Thus, for images with less than  $\frac{2}{3}N^{\frac{2}{3}}$  pixels, image-based rendering will be faster than the parallel visualization server. It is interesting to note that this expression is independent of both the number of isosurfaces and the network bandwidth and latency.

**Conclusion 1:** If image resolution is fixed, image-based rendering becomes increasingly attractive compared to parallel visualization servers as the data set size increases.

We analyze the performance of the subsampling techniques by finding the value of the reduced data set size,  $N_R$ , reported as a percentage of N,  $\mathcal{P}_N = \frac{N_R}{N} \cdot 100$ , such that the costs of the subsampling model are equal to the smaller of either Model 1 or 2 costs. For both uniform and multiresolution grid subsampling, we start with a representative set of base parameters  $N = 128^3$ , P = 8, R = 10, and I = 64. To examine the effect of these parameters on  $\mathcal{P}_N$ , we vary each one individually while holding the others fixed, and plot the results in Figure 1. Values of  $\mathcal{P}_N$  below the curves indicate the regimes for which subsampling is more cost-effective than either parallel visualization servers or image-based rendering at a resolution of  $X = 512^2$ .

As expected, uniform grid subsampling can use a higher percentage of grid points than can be used with multiresolution grid subsampling. In fact, for this base problem size it is often the most cost-effective strategy for remote visualization. In contrast, the multiresolution subsampling is cost-effective when  $\mathcal{P}_N < 12$ . However, the multiresolution approach can cluster points in regions of interest so that fewer grid points are needed to achieve the same fidelity to the original data set. These tradeoffs are examined in more detail in Section 4.3. In general, the following

Method	$N_R$	$\mathcal{P}_N$	Avg $\sigma_n$	Max $e_n$	]	Method	$N_R$	$\mathcal{P}_N$	Avg $\sigma_n$
Uniform	4096	7.5	.0652	.932		Uniform	202799	9.9	1.08
Uniform	7179	14.0	.0506	.763		Uniform	474443	23.3	.701
Uniform	15616	28.9	.0347	.569		Uniform	800602	39.1	.699
Uniform	23560	43.6	.0230	.555		Uniform	1212327	59.2	.268
	Criter	ia = Avg	g. $\sigma_n$		Criteria = Avg. $\sigma_n$				
Multi-Res	808	1.5	.114	1.24		Multi-Res	62642	3.1	.568
Multi-Res	1630	3.0	.0703	1.24		Multi-Res	124667	6.1	.343
Multi-Res	3268	6.0	.0397	1.08		Multi-Res	250790	12.2	.176
Multi-Res	6550	12.1	.0238	1.05		Multi-Res	502772	24.5	.080
Multi-Res	13108	24.2	.0083	.515					

Table 2: Results for uniform grid and multiresolution data reduction techniques<br/>2D Rayleigh Taylor3D Hairpin Vortices

trends are evident:

**Conclusion 2:** The network speed is not the primary bottleneck in any of these strategies as varying the value of the parameter R has a minimal effect on the results.

**Conclusion 3:** Subsampling approaches are best used for moderate problems sizes and for problems with a large number of visualization tasks. As the problem size increases, the size of reduced data set that gives good performance relative to the other two strategies decreases causing larger approximation errors. We note that this effect can be somewhat mitigated by the use of a windowing functionality that allows the user to obtain higher resolution only in regions of interest.

#### 4.3 Uniform and Multiresolution Subsampling

For a given set of parameters, the graphs in Figure 1 clearly show that uniform grid resampling is significantly more cost-effective than multiresolution subsampling. To determine relative performance benefits, we computed the ratios of the uniform and multiresolution subsampling breakeven percentages. In most cases, uniform subsampling is about a factor of 12 more cost-effective than multiresolution subsampling. The primary difference between the two methods is the large difference in the cost of generating isosurfaces,  $C_{I_u}$  and  $C_{I_m}$ . In fact, if we consider a case in which  $C_{I_m}$  is  $5.0 \cdot 10^{-6}$ , or about four times slower than  $C_{I_w}$ , the corresponding performance ratio is 4.

Thus, the multiresolution technique will outperform the uniform grid method only if the subsampling approximation errors are reduced by the same amount or more using significantly fewer grid points. To explore these tradeoffs, we subsample two different application data sets. The first data set is from a two-dimensional Rayleigh-Taylor (R-T) simulation and contains  $N = 5.3 \cdot 10^4$ data points. This data set is characterized by a contact discontinuity between two fluids of differing density and represents a broad class of applications whose primary features are sharp discontinuities which are typically local, lower-dimensional phenomena. The second data set is from a threedimensional simulation of hairpin vortices developing in flow around a hemisphere and contains  $N = 2.05 \cdot 10^6$  data points. This problem is representative of a class of applications in which the scalar field of interest describes full-dimensional features.

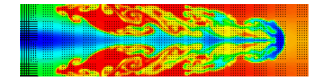
For each problem, we create reduced data sets of approximately 5, 10, 25, and 50 percent in

the uniform grid case and approximately 1.5, 3, 6, 12, and 25 percent in the multiresolution case. For multiresolution subsampling,  $N_R$  was specified, and the code automatically determined the best decomposition to minimize the standard deviation,  $\sigma_n$ . In Table 2, we report the the average  $\sigma_n$  and maximum  $e_n$  over all octants. The first value gives a measure of the overall fidelity of the reduced data set to the original data set; the latter value gives a worst-case measure of fidelity. For the hairpin vortex data set, the scalar field is interesting in that only values less than negative one are of interest; all other values are disregarded as noise. These noise values can vary dramatically outside the regime of interest, rendering the maximum deviation error measure ineffective.

In both cases, we achieve the same average errors using far fewer multiresolution grid points than uniform grid points. For the R-T problem, the same average error is achieved by approximately a factor of five fewer grid points; for the hairpin vortex data set, the factor is ten.

In the left image in Figure 2, we show the R-T data set subsampled using a uniform grid containing 15616 points. On the right, we show the same data set subsampled with the multiresolution technique using 4102 points; points are clustered at the discontinuity. The average errors are .0347 and .0339, respectively. In Figure 3, we show isosurfaces from the hairpin vortex set for vorticity indicator of -.28. The left figure shows uniform grid subsampling using 34798 points which results in an average error of 3.18. The right figure shows multiresolution subsampling with 34725 grid points which results in an average error of 1.46.

**Conclusion 4:** For the cost parameters  $C_{I_u}$  and  $C_{I_m}$  considered here, uniform grid subsampling outperforms multiresolution subsampling when the average error criteria is used. As techniques for generating multiresolution isosurfaces improve in speed, multiresolution subsampling approaches will become increasingly cost-effective.



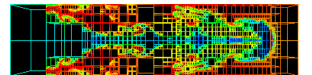


Figure 2: The R-T problem subsampled by uniform (left, 15616 points) and multiresolution (right, 4102 points)

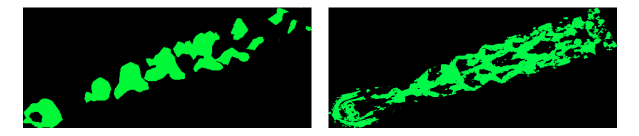


Figure 3: The hairpin vortex data set subsampled by uniform (left, 34798 points) and multiresolution (right, 34725 points) methods.

## **5** Conclusions

From our analysis of the computation and communication costs associated with three different strategies for remote, interactive exploration of large data sets, we find that each has regimes for which is it is the most cost-effective approach. In general, the subsampling approaches will be the most cost-effective for moderate problems sizes containing tens of millions of elements, followed by parallel visualization servers and image-based rendering as the problem size increases.

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