TLS and all that

Ethan A Merritt

CCP4 Summer School 2009 (Argonne, IL)

Abstract

TLS (Translation/Libration/Screw) is a mathematical model that predicts the local positional displacement of atoms in a crystal structure based on an underlying assumption that each atom acts as a member of a rigid body that is displaced normally about a mean position. The net displacement contains contributions from actual vibration of the molecule in the crystal and from the static disorder within the crystal lattice that results when different unit cells trap different microconformers of the molecule as it exists in solution. The averaged displacement of each atom is seen crystallographically as nonspherical electron density at the atomic positions. The TLS formalism was originally developed in the late 1960s to help interpret the 'thermal ellipsoids' used in small molecule crystallography at high resolution [Schomaker & Trueblood, 1968, Dunitz et al., 1988]. Thus it was a method of analysis rather than of refinement. Its success in identifying rigid groups by examination of the individual atomic displacement parameters in small molecule structures makes it attractive for use also in macromolecular crystallography, where it can be used to find domain and loop flexibility in proteins. Moreover, the equations that derive a single 20-parameter TLS description from the observed thermal ellipsoids of many atoms can be turned around to great benefit. In this reverse direction form, the TLS description is used to predict the individual atomic displacements of many atoms. This turns out to be a great way to describe and refine protein structures. Protein crystals rarely yield the very high-resolution data (≤ 1.2 Å) that would be needed to model anisotropy (thermal ellipsoids) for every atom separately. But even at low resolution (>3Å) they yield sufficient data to refine TLS models, and these TLS models describe aspects of the crystalline protein that would otherwise not be described at all: dynamic and static disorder due to flexibility¹.

¹A popular corollary is that better descriptions of the crystal content allow improved values of F_{calc} , and therefore TLS models generally allow better R factors.

A choice of ways to describe the displacement of individual atoms

Consider a crystal of the hypothetical protein Longcase. The crystal lattice is established by tight packing of the exterior "Case" domain. The separate "Pendulum" domain is relatively unconstrained by the lattice, and hence is present in a mixture of conformations. Crystallographically we will observe an average of these multiple conformations, which will lead to slight blurring at the hinge-point of this domain but very substantial blurring of the large atom at the bottom of the domain. How best to model this?

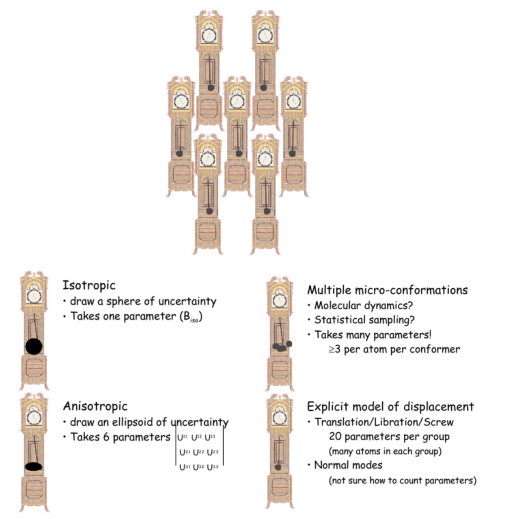


Figure 1: Modeling displacement due to an obvious vibrational mode

If Longcase crystals diffract to atomic resolution, we may have the option of modeling the obviously anisotropic displacement of the pendulum tip as an ellipsoid. At lower resolution we don't have that choice. But even at such high resolution it may be worth using an explicit model of the vibrational mode that causes this anisotropy. We will consider only TLS models, although other descriptions are possible.

Simple models vs. complex models - how many TLS groups?

In the Longcase example it was obvious that the structure can be divided into two parts, only one of which is moving/vibrating. It is obvious which part that is, and that it really is acting as a rigid body. In the general case we do not know how many flexible parts there are, or how finely we would have to slice up the structure to reach a point where the motion of each individual part is well approximated as a rigid body.



Figure 2: Modeling the swing of a bat

In choosing how many separate pieces should be modeled, it also helps to have a question in mind. To ask "what part of the baseball player may contact the ball?", we only need a simple model. Perhaps it is sufficient to consider arms, bat, and body. On the other hand, if we want to ask "are the ankles important to the swing?", a more complex model is needed.

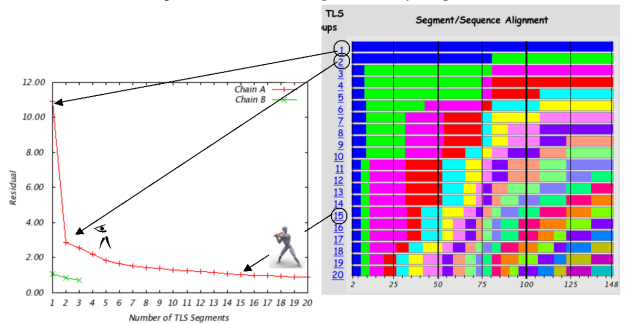


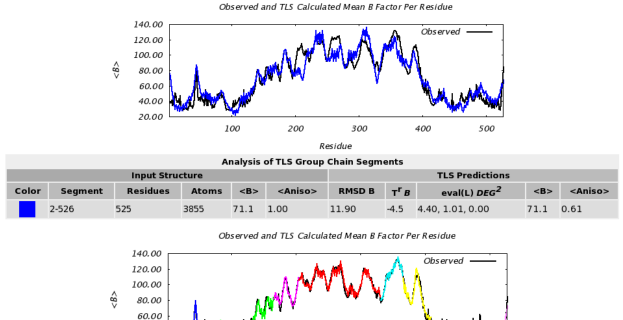
Figure 3: TLSMD: Modeling the flexibility of a protein

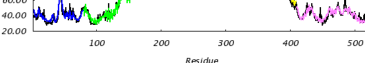
Overview of TLS refinement guided by TLSMD

- 1. [incredibly important!] start with a refined set of B values
 - usual case: conventional B_{iso} refinement
 - high resolution: anisotropic U^{ij} refinement
 - low resolution: individual B_{iso} or group B per-residue
 - DON'T use the original Bs from a molecular replacement solution
 - DON'T use fixed B. Models with constant B are useless; newly-built residues often have B=15.
- 2. [*the easy part*] submit your starting model to the TLSMD server http://skuld.bmsc.washington.edu/~tlsmd
- 3. [*the server does the hardest part*] generate optimal multi-segment TLS models that best explain the observed B values
- 4. Choose the TLS model you prefer; i.e., how many segments per chain?
 - The eventual R value is expected to track the change in residual reported by the TLSMD server. If there isn't much drop in residual as you add TLS groups, probably TLS is not going to help your R factors much. My rule of thumb is that if the residual doesn't eventually drop by a factor of 2, a multi-group TLS model is only worth pursuing if you care about the hinge points or domain boundaries for their own sake.
 - Look for a dog-leg in the plot. For example, Figure 3 shows a huge dog-leg at 2 segments and a less dramatic one at 5 segments.
 - Look down the list of partitions for stable locations of implied hinge-points.
 - The fit of observed to predicted B should be equally good for all segments (Figure 4). But if it isn't, this may indicate a problem in the original refinement rather than a problem with TLS analysis per se.
 - Does the model make sense? The animations may help you decide.
- 5. [*Optional*] simplify the model further (Figure 6).
- 6. [refmac or phenix does this part] refine the TLS model against the crystallographic data
 - You must choose whether to reset the starting B_{iso} values to a constant. If in doubt, try it both ways and see which works better (Figure 5).
- 7. [... and this part] refine the TLS model jointly with coordinates and maybe residual contribution Biso
 - High to medium resolution: refine residual B_{iso} for each atom (Figure 5a).

- Medium to low resolution: refine only an overall B, *not* individual B_{iso} for each atom. This gives you a pure TLS model (Figure 5b).
- 8. repeat steps 2-7. Remember, the quality of the multi-group TLS model suggested by TLSMD is limited by the quality of the B values in the PDB file you give it for analysis. As your model improves, the TLS description may improve also (Figure).
- 9. Figure out how to tell the PDB what you have done. Unfortunately, there is still a great deal of confusion about what exactly is in a PDB file that has been refined with TLS. Many programs ignore it. For example, if you deposit a model refined with TLS, the validation program run by the PDB (sfcheck) usually flags your structure factor file as having poor correlation with your model. In other words, it cannot confirm your F_{calc} values because it fails to apply the TLS model even though it is listed in your PDB file.

Figure 4: A good choice of TLS segmentation will give roughly equal residuals for each segment. The curves below show the average per-residue agreement of a 1-group TLS model (top) and a 7-group TLS model (bottom) for the same protein structure. It is clear that some portions of the chain are not well described by the single TLS group model. However, after partitioning the chain into 7 separate TLS groups, the fit is of roughly equal quality everywhere.





Resid

Analysis of TLS Group Chain Segments

Input Structure						TLS Predictions				
Color	Segment	Residues	Atoms		<aniso></aniso>	RMSD B	т ^г в	eval(L) <i>DEG</i> ²		<aniso></aniso>
	2-78	77	569	41.0	1.00	5.73	32.9	10.58, 0.05, 0.00	41.0	0.80
	79-164	86	597	51.9	1.00	5.25	28.7	12.46, 0.00, 4.03	51.9	0.66
	165-203	39	297	87.9	1.00	4.35	72.4	8.18, 3.41, 0.00	87.9	0.86
	204-329	126	942	105.0	1.00	5.05	52.3	5.63, 1.34, 0.00	105.0	0.87
	330-364	35	272	113.6	1.00	3.18	110.2	4.75, 0.00, 0.00	113.6	0.86
	365-408	44	324	84.7	1.00	6.07	70.3	9.42, 0.01, 0.00	84.7	0.77
	409-526	118	854	42.6	1.00	5.71	30.0	5.36, 2.26, 0.00	42.6	0.82

Run Refmac5 Initial parameters from /sgpp/data/pil soaks/1554b3/ccp4/CCP4 DATABASE/	7 refmac5	def - X	Run Refmac5 Initial parameters from /sgpp/data/pil soaks/1554b3/ccp4/CCP4 DATABASE/	7 refmac5	def - X
		Help			Help
Job title TLS plus individual Biso refinement, Reset residual B factors to a constant before refining	Job title Pure TLS model, NO RESIDUAL Biso				
Bo TLS & restrained refinement _ using no prior phase information _ input		po TLS & restrained refinement using no prior phase information input			
no twin refinement			no - twin refinement		
MTZ in 1554b3 - 1554b3_truncate1.mtz	Browse	View	MTZ in 1554b3 - 1554b3_truncate1.mtz	Browse	View
FP F_1554b3 Sigma SIGF_1554b3			FP F_1554b3 Sigma SiGF_1554b3		
MTZ out 1554b3 - 1554b3_refmac04TLSI.mtz	Browse	View	MTZ out 1554b3 - 1554b3_refmac04TLSI.mtz	Browse	View
PDB in 1554b3 - 1554b3_refmac03.pdb	Browse	View	PDB in 1554b3 1554b3_refmac03.pdb	Browse	View
PDB out 1554b3 - 1554b3 refmac04TLSLpdb	Browse	View	PDB out 1554b3 1554b3_refmac04TLSLpdb	Browse	View
LIB in 1554b3 - Merge LIBINS	Browse	View	LIB in 1554b3 - Merge LIBINS	Browse	View
Output lib 1554b3 - 1554b3.cif	Browse	View	Output lib 1554b3 — 1554b3.cif	Browse	View
TLS in (optional) 1554b3 - 8groupTLS.tlsin Create TLSIN	Browse	View	TLS in (optional) 1554b3 - 8groupTLS.tisin Create TLSIN	Browse	View
TLS out 1554b3 - 1554b3_refmac04TLSLtis			TLS out 1554b3 - 1554b3_refmac04TLSI.tls	Browse	View
Include keyword file 1554b3 - Browse View			Include keyword file 1554b3 -	Browse	View
a Harvesting Data Harvesting					
TLS Parameters			TLS Parameters		~
Number of cycles of TLS refinement 15			Number of cycles of TLS refinement 15		
Set initial Bfactors to 20.0 (numeric value unimportant) Your choice whether to keep curren	tBorn	ют	Set initial Bfactors to 20.0 (numeric value unimportant)		
Refiniement Parameters Refiniement Parameters					
Do 10 cycles of maximum likelihood restrained refinement after TLS refinement			Do 10 cycles of maximum likelihood restrained refinement after TLS refinement		
Use hydrogen atoms: generate all hydrogens — and 🗌 output to coordinate file		-	Use hydrogen atoms: generate all hydrogens and output to coordinate file		-
Resolution range from minimum 59.190 to 1.860					
domatic weighting 🗢 Use experimental sigmas to weight Xray terms					
genergifting term (0.08					
e isotropic temperature factors					
Exclude data with freeR label ErceR_flag with value of 0					
Run 🛁 Save or Restore 🛁	Close		Run — Save or Restore —	Close	

Figure 5: Telling refmac whether to refine a residual B_{iso} for each atom (a), or a pure TLS model (b)

Figure 6: **Optional Step: Merge discontiguous chain segments into a single TLS group.** Domains are often formed from segments of the polypeptide chain that are not adjacent in the sequence. A simple example of this is shown below in which a 5-strand beta sheet is formed by the N- and C- termini. In a case like this we expect that the whole domain may act as a single approximately rigid body, and be well described as a single TLS group. The automated analysis is not currently smart enough to recognize this by itself, although it does print out tables of how well the individual TLS descriptions for each segment agree with each other. The diagonal terms in the table are the residuals showing how well the TLS model for each segment predicts the B values reported for that segment. The off-diagonal terms are the residuals you would get after combining two groups into a single group. If a pair of off-diagonal terms are approximately the same magnitude as the on-diagonal terms for those same segments, this is a hint that they may actually act as a single group.

RMSD B Values of Combined TLS Groups

	2-138	139-395	<mark>396-526</mark>
2-138	5.98	12.25	7.08
139-395	12.25	6.24	10.54
<mark>396-526</mark>	7.08	10.54	6.19

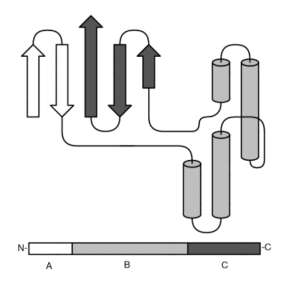
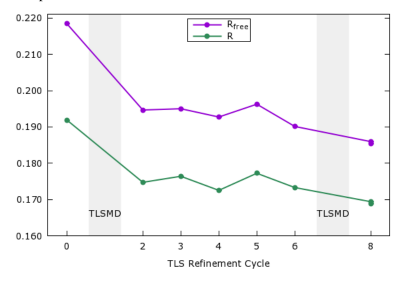


Figure 7: **Revisiting the TLSMD model after refinement.** You should probably re-evaluate your TLS model as your refinement nears completion. TLSMD constructs a model that optimally predicts the distribution of B values in the 3-dimensional space occupied by your structure. After you have refined the structure you hopefully have better B values and 3D coordinates, so TLSMD analysis may be able to construct a better TLS description.



Case study: TLS Refinement at 3 Å resolution

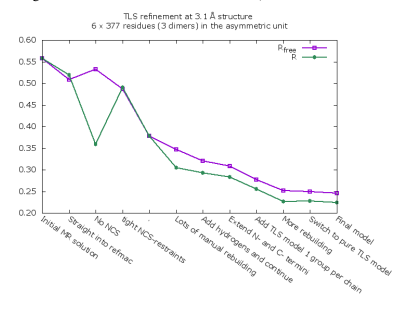


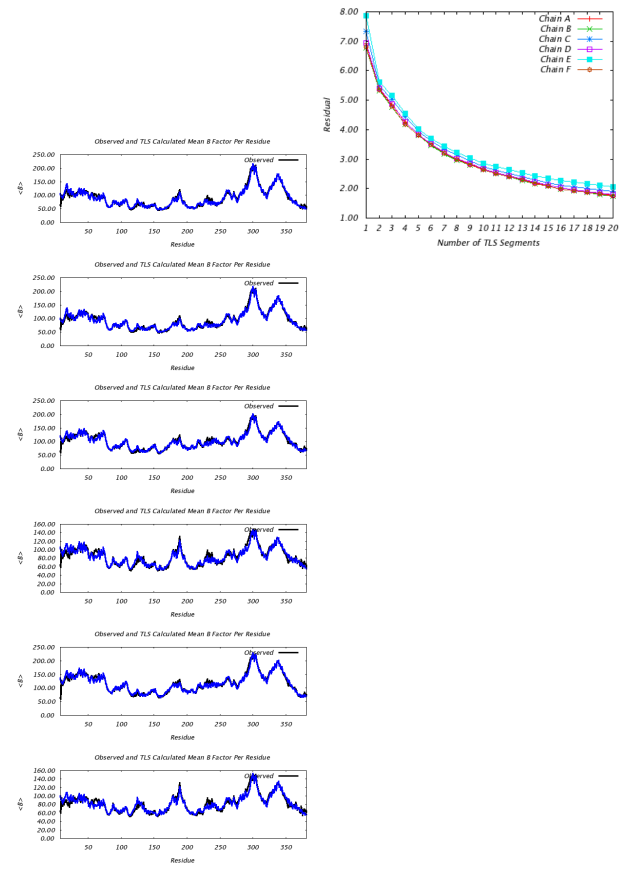
Figure 8: Refinement at 3 Å resolution, from initial MR to final model

Here is the course of refinement of a real protein at 3 Å resolution. Phaser was able to place 3 copies of a homologous dimer in the asymmetric unit. Conventional refinement of the initial model (individual B_{iso} , no special restraints) was not well-behaved. Adding tight NCS restraints helped a lot. The usual iteration of manual rebuilding with refinement of an NCS-restrained model using individual (NCS-restrained) B_{iso} stalled out at $R = 0.28/R_{free}=0.31$. At this point I submitted the model to TLSMD. The analysis is shown in Figure 9. Several things are apparent. Partitioning the chain into many segments yields an overall drop in the residual by about a factor of 4. That is pretty good.

However, it is interesting that even a single group TLS model fits the refined B_{iso} very well (Figure 9). This suggested that it was worth refining a very simple, one group per chain, TLS model first. Indeed, adding this simple TLS model to the refinement lowered the crystallographic residuals to $R = 0.23/R_{free} = 0.25$. That is pretty dramatic, and hints that the primary source of displacement in the crystal is the rocking of entire molecules within the lattice (remember that this TLS in this case is treated an entire monomer as a rigid body). This hint is borne out when the refinement is switched over to using a pure TLS model, still treating each monomer being as a rigid body, with no individual B_{iso} terms. This even simpler model yields slightly better *R* and R_{free} than the model which included individual *B* terms.

Figure 9: TLSMD analysis of the conventionally refined model in case study #1.

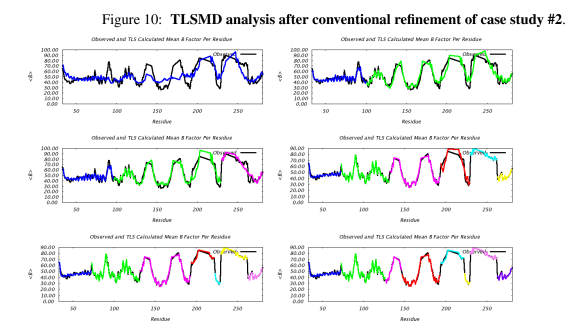
(a) 1-group TLS fits for each chain A B C D E F (b) Overall residual as a function of number of TLS groups



For whatever reason, in this case partitioning each chain into many segments does not significantly drop *R* or R_{free} beyond this point. Refining a TLS model with 12 groups per chain yielded $R = 0.225/R_{free} = 0.252$. This seems contrary to what I would normally predict based on the curve in Figure 9, but it is consistent with the observation that a 1 group per chain model already fits very well everywhere (Figure 9). Compare this, for instance, to the more typical single group curve in Figure 4. So for this particular crystalline protein, TLS refinement was extremely effective even without splitting the chains into multiple TLS groups.

Case study #2: TLS Refinement at 2.8 Å resolution

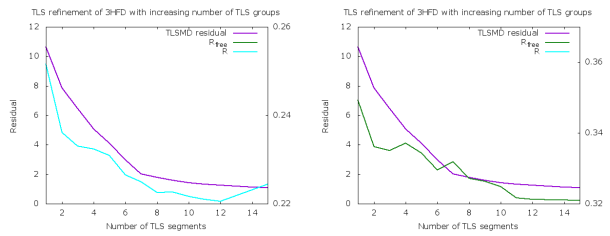
Here is another example of refinement at moderately low resolution, one that works out very differently. In this case the protein is a dimer, but the asymmetric unit contains a single monomer of 270 residues. Again the initial structure solution came from molecular replacement. Manual rebuilding and conventional B_{iso} refinement brought yielded $R = 0.249/R_{free} = 0.341$.² TLSMD analysis of the model indicated a distinct dog-leg in the residual plot at 7 groups. You can see essentially the same thing by noticing that the perresidue fit of predicted to observed B factors is very uneven for partitions with fewer segments (Figure).



In this case these features of the TLSMD analysis do indeed correlate with the improvement in R and R_{free} obtained from refinement of TLS models with an increasing number of TLS groups (Figure 11).

 $^{{}^{2}}R$ and R_{free} are rather far apart, which probably indicates model bias. Unfortunately in this case we had no NCS or experimental phases to help overcome this bias.

Figure 11: TLS refinement after TLSMD analysis of case study #2. These are pure TLS models, i.e. no B_{iso} terms are included in the model.



Further reading

Theory of TLS

[Rosenfield et al., 1978]

[Willis & Pryor, 1975]

TLSMD server

[Painter & Merritt, 2006a]

[Painter & Merritt, 2006b]

References

[Dunitz et al., 1988] Dunitz, J., Schomaker, V., & Trueblood, K. N. (1988). J. Phys. Chem. 92, 856-867.

[Painter & Merritt, 2006a] Painter, J. & Merritt, E. A. (2006a). Acta Cryst. D62, 439-450.

[Painter & Merritt, 2006b] Painter, J. & Merritt, E. A. (2006b). J. Appl. Cryst. 39, 109–111.

[Rosenfield et al., 1978] Rosenfield, R. E., Trueblood, K. N., & Dunitz, J. D. (1978). Acta Cryst. A34, 828–829.

[Schomaker & Trueblood, 1968] Schomaker, V. & Trueblood, K. N. (1968). Acta Cryst. B24, 63-76.

[Willis & Pryor, 1975] Willis, B. T. M. & Pryor, A. W. (1975). *Thermal Vibrations in Crystallography*. Cambridge University Press.

Acknowledgments

Source for complex model baseball swing images: http://blogs.tamu.edu/ariel [Ariel Chisholm, Texas A&M University Visualization Sciences Coursework]