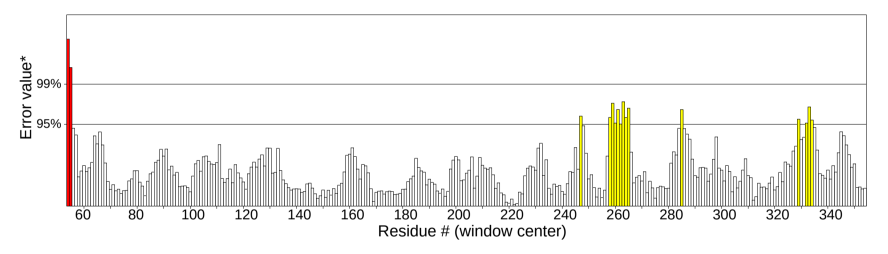
Program: ERRAT2 File: .CYP1B1.pdb Chain#:A

Overall quality factor**: 94.595

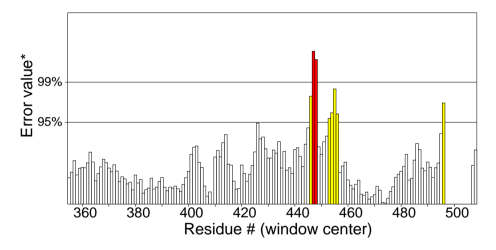


*On the error axis, two lines are drawn to indicate the confidence with which it is possible to reject regions that exceed that error value.

^{**}Expressed as the percentage of the protein for which the calculated error value falls below the 95% rejection limit. Good high resolution structures generally produce values around 95% or higher. For lower resolutions (2.5 to 3A) the average overall quality factor is around 91%.

Program: ERRAT2 File: .CYP1B1.pdb Chain#:A

Overall quality factor**: 94.595



*On the error axis, two lines are drawn to indicate the confidence with which it is possible to reject regions that exceed that error value.

^{**}Expressed as the percentage of the protein for which the calculated error value falls below the 95% rejection limit. Good high resolution structures generally produce values around 95% or higher. For lower resolutions (2.5 to 3A) the average overall quality factor is around 91%.