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Subject: Re: New DPMgen results disagree with earlier simulations

Posted by [Stefano Spataro](#) on Fri, 05 Feb 2010 16:07:36 GMT

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However,

PndDpmDirect does not propagate anymore this number, which is not so nice and must be fixed.

I am wondering which could be a good value for thetamin, and how much this can change the final results for elastic coulomb, to use it as default value instead of zero.

Do you have some idea on what could be used as default parameter?

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