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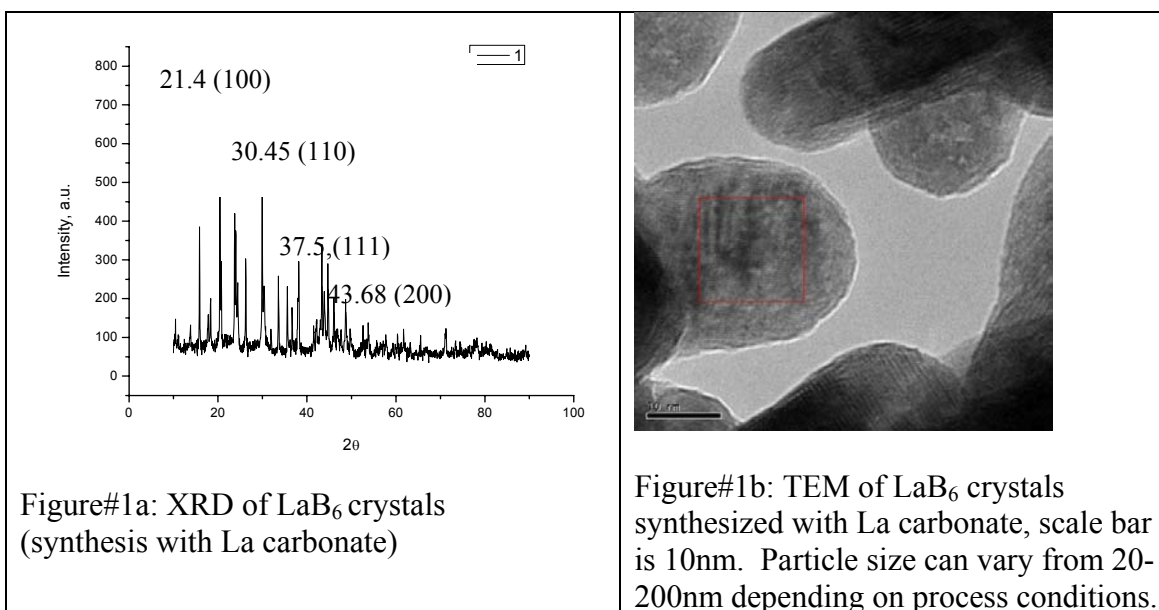
New Synthesis for Lanthanum Hexaboride Nanocrystals

Lisa Pfefferle

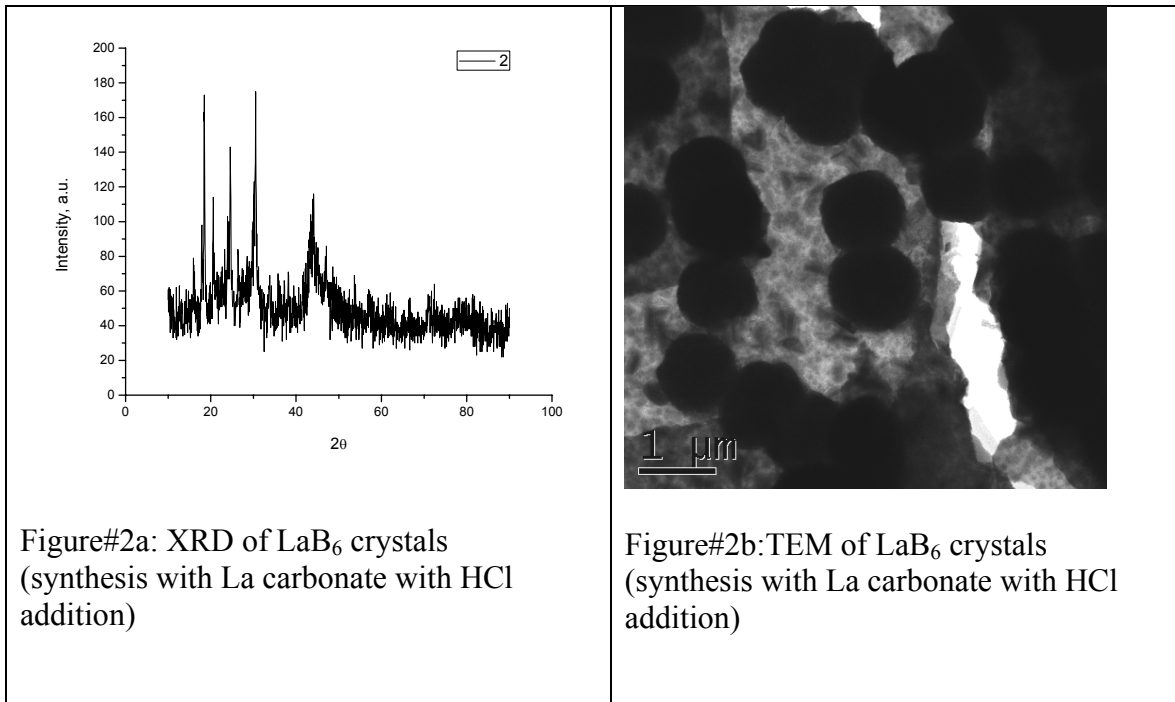
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Although lanthanum hexaboride has been used for many years as a cathode material, new applications exploiting its unusual electron emission properties are now being investigated [Boustani *et al.*, Monnier and Delley]. Nanocrystalline LaB₆ materials have been predicted to provide important advantages for IR absorbers as well as for nanoelectronics in general because of their low work function. Lanthanum hexaboride is usually made by mixing La and B with Al in a furnace at 1500K. The molten flux method precipitates large crystals (see *eg.* Inoue *et al.*). This method, however, is not suitable for *nanocrystal* production.

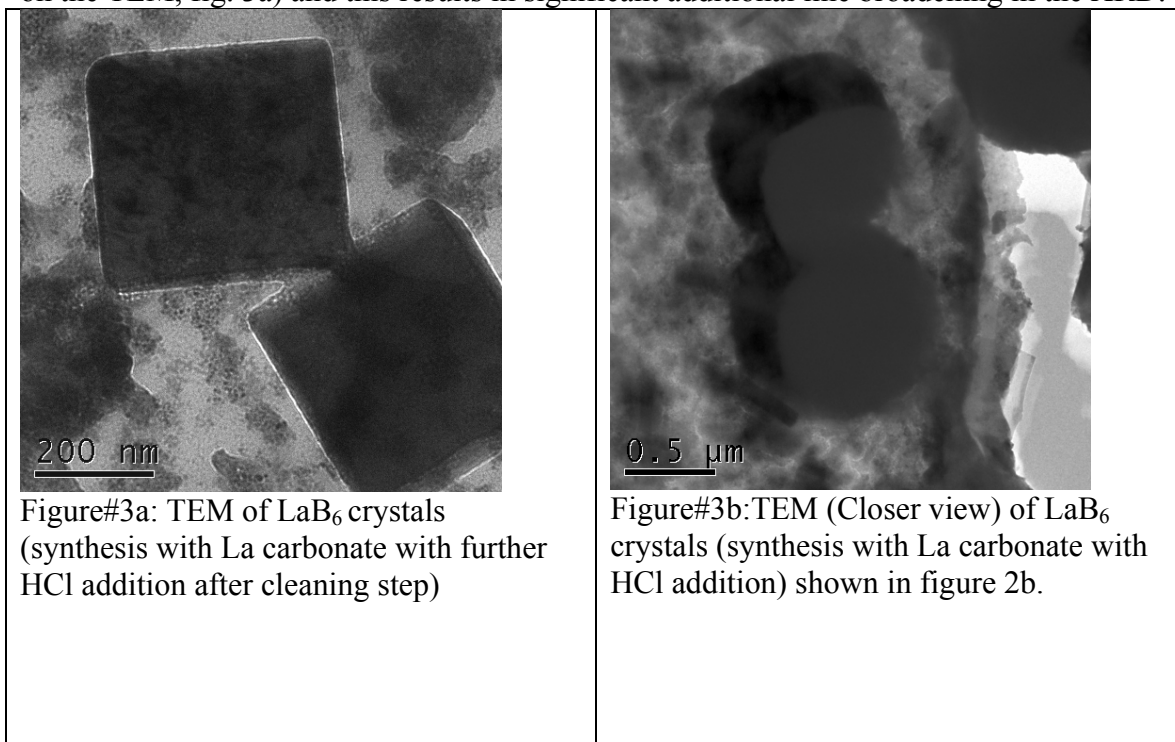
We have synthesized LaB₆ nanocrystals for the first time using a low temperature sonochemical synthesis. Several different types of crystals were synthesized. First, 20-200nm diameter crystals were synthesized using lithium borohydride and lanthanum carbonate sonicated in a tetrahydrofuran solvent. These showed a good XRD pattern confirming the lanthanum hexaboride structure (see figure #1a and b below).



We also showed that we can control the thickness of the crystals using addition of HCl during the synthesis. A small amount of HCl acid in the synthesis leads to thinner/smaller crystalline domains and reduces the amorphous material; adding more HCl results in further thinning as observed by further significant broadening of the peaks in XRD. (see XRD Figure #2a where as acid is added the lines widen considerably).



The crystals in one run adding additional HCl were very thin (you can see through them on the TEM, fig. 3a) and this results in significant additional line broadening in the XRD.



A different synthesis strategy using Lanthanum chloride as the La precursor resulted in smaller (several nm diameter) boron/lanthanum/oxygen particles which could likely be annealed at high T to give LaB₆. Another strategy using the same precursor with water as the solvent resulted in a purple powder containing amorphous boron/lanthanum particles 3-5 nm in diameter which will likely be able to be converted to LaB₆ on annealing at high temperature. An oven is currently being installed that will allow us to anneal samples in varied environments up to 1900K.

Thus several different routes to varied structure LaB₆ nanocrystals have been demonstrated. This is extremely promising because it means that the LaB₆ crystals can be designed by varying the synthesis strategy to meet varied application needs. Initial work suggests that reasonably narrow diameter distributions of crystals with mean diameters ranging from 5-200nm can likely be synthesized and that the aspect ratio can be varied and surface orientation can likely be varied. Issues that need to be addressed in future research are parameter optimization to control particle structure and purity issues.

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