

Avoiding adsorption of DNA to polypropylene tubes and denaturation of short DNA fragments

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▼ **Two problems can arise when working with small quantities of DNA in polypropylene tubes: first, significant amounts of DNA can become lost by sticking to the tube walls; second, short DNA fragments tend to denature when binding to polypropylene. In addition, DNA also tends to denature upon dehydration. We have found that a simple way to solve these problems is by using polyallomer tubes instead of polypropylene and by avoiding certain salts, such as sodium acetate, when drying DNA.**

DNA is usually stored in polypropylene tubes, which have become widely used for their resistance to solvents, their strength, their ease of use and their low price. Polypropylene is a very hydrophobic material, whereas DNA is a highly charged macromolecule, two characteristics that minimize the interactions of DNA with tube walls and tend to avoid the adsorption problems often found with other macromolecules, especially proteins.

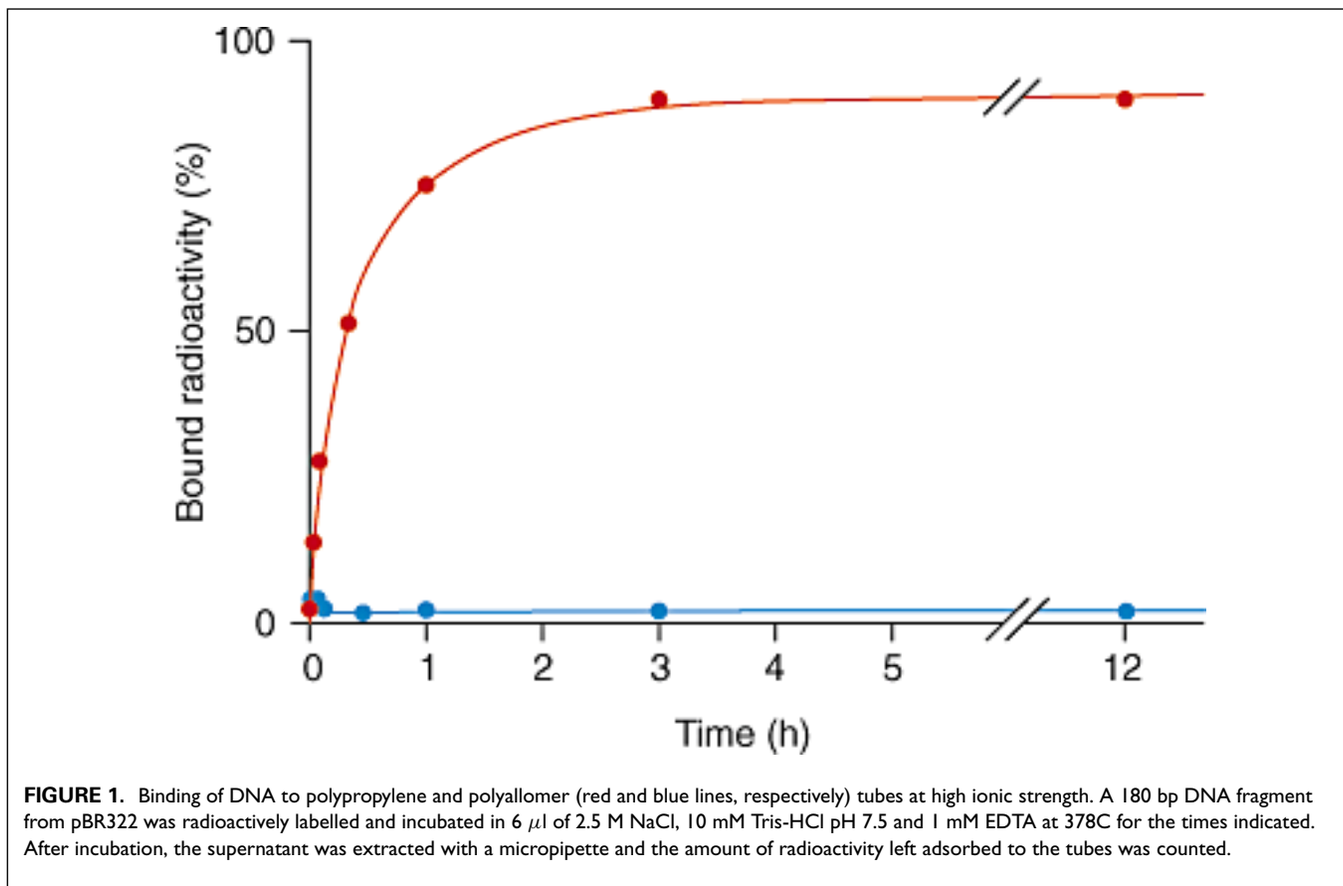
But DNA can, in fact, bind to polypropylene tubes. This is particularly striking at high ionic strength: Fig. 1 shows that DNA fragments bind quickly to polypropylene tube walls in 2.5 M NaCl, with 75% of the material adsorbed after 1 h and 90% after 3 h in this experiment. Under such conditions, the amount of adsorbed DNA can be as high as 5 ng/mm² of tube wall. Tests performed at ionic conditions varying from TE (10 mM Tris-HCl, 1 mM EDTA, pH 7.5) to TE plus 0.5 M NaCl, showed that DNA also sticks to tube walls at lower ionic strength, but with important variations between different batches of tubes. Tests were performed

with 1 ng of radioactively labelled DNA per tube in 10 μ l. With some batches of tubes, the percentage of adsorption was always high (between 80% and 95%; such tubes actually had a binding capacity as high as >10 ng for a volume of 10 μ l). With other tube batches from the same manufacturer (Eppendorf) the percentage of adsorption varied apparently at random between 5% to 95%, all conditions being kept identical. Finally, other batches always presented little adsorption (<10%) at low ionic strength. In all those tests, using single-stranded instead of double-stranded DNA made no significant difference.

The problems resulting from this adsorption, loss of material and denaturation of adsorbed fragments (see below), led us to search for different kinds of tubes in which DNA would not adsorb to the tube walls. Upon testing the three different kinds of 1.5 ml plastic tubes sold by Beckman, which are made of polypropylene, polyethylene, or polyallomer, respectively, we observed the same adsorption of DNA to polypropylene as before, and a similar adsorption to polyethylene. In strong contrast, no adsorption of DNA to polyallomer tubes was observed (Fig. 1), irrespective of the ionic strength. We have used polyallomer tubes for DNA storage since this finding, and although such tubes are reported to be slightly less resistant to aromatic and halogenated hydrocarbons than polypropylene tubes, the difference is not obvious and we have had no technical problems using them. Their only disadvantage is their price, which is currently about ten times higher than that of polypropylene tubes.

Another problem with polypropylene tubes comes from their tendency to denature DNA fragments. Storing short DNA fragments in polypropylene tubes often induces a

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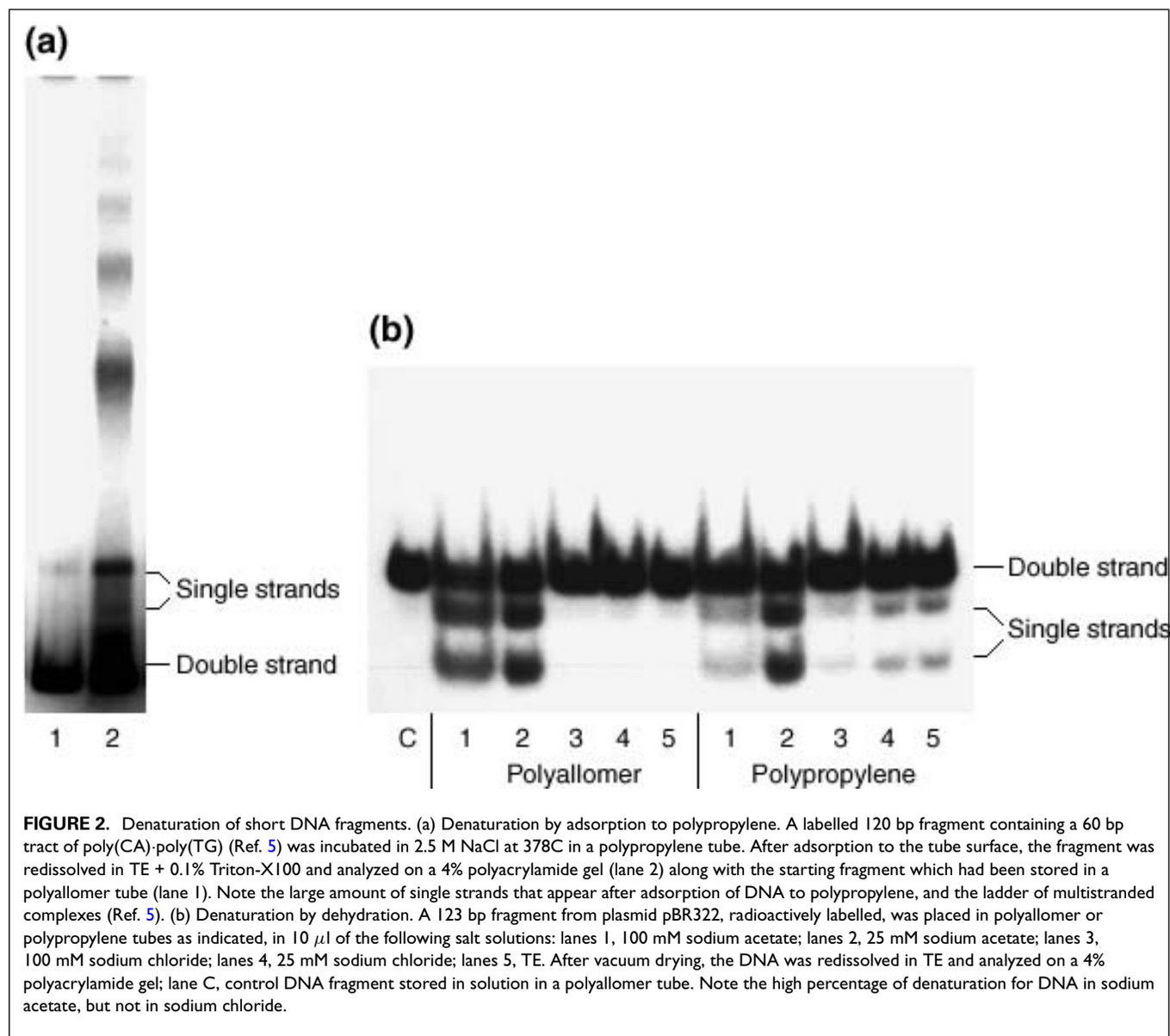
significant amount of strand separation, a phenomenon that has been well documented by Belotserkovskii and Johnston (Ref. 1, 2, 3), and also observed by us (Ref. 4). Instead of stabilizing the double helix as expected, the high ionic strength stimulates the strand dissociation as well as the interaction of DNA with polypropylene, as shown in Fig. 2. Incidentally, such conditions of high salt concentration also lead to the formation of very interesting multi-stranded complexes, a detailed study of which will be presented elsewhere (Ref. 3, 5; Gaillard *et al.*, unpublished; see the ladder of bands in Fig. 2a, lane 1).

As in the case of adsorption of DNA to tube walls, we have found that using polyallomer tubes is the most simple and straightforward way to avoid this DNA denaturation.

Svaren and Chalkley (Ref. 6) also showed the tendency of short DNA fragments to denature upon dehydration, in particular during the dehydration step that usually follows ethanol precipitation. We have observed that this dehydration-induced denaturation depends strongly on the nature of the salt. Figure 2b shows the same DNA fragment dissolved in two different salt solutions, sodium chloride and sodium acetate, and vacuum dried. No denaturation is observed with the fragment in sodium chloride, whereas an

important percentage of denaturation is observed with the fragment in sodium acetate. We do not know the explanation for this difference. The precipitation efficiency being identical, we now use sodium chloride instead of sodium acetate for ethanol precipitation of DNA. Also, note again on Fig. 2b the superiority of polyallomer tubes, compared with polypropylene, for storage of DNA in its native state.

We now always use polyallomer tubes for DNA storage and strongly recommend them for storage of small amounts of short DNA fragments. On the other hand, it is possible to prevent DNA from interacting with polypropylene, the best way being by addition of 0.1% of a non-ionic detergent such as Triton X-100. Siliconizing tubes is an effective measure against DNA adsorption, but does not entirely suppress DNA denaturation on tube walls. Some manufacturers sell 'low-binding' tubes; indeed, we observed little adsorption of DNA with the ones we tested. However, as long as the exact nature of these tubes is kept a secret by manufacturers, the risk exists that some chemical added to polypropylene or to the tube surfaces will contaminate DNA. Examples can be found in the literature where a contamination by a substance released from the tubes has been well documented (Ref. 3, 7). In addition, we do not know whether DNA binds



to polypropylene itself, or to some minor component of the plastic.

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