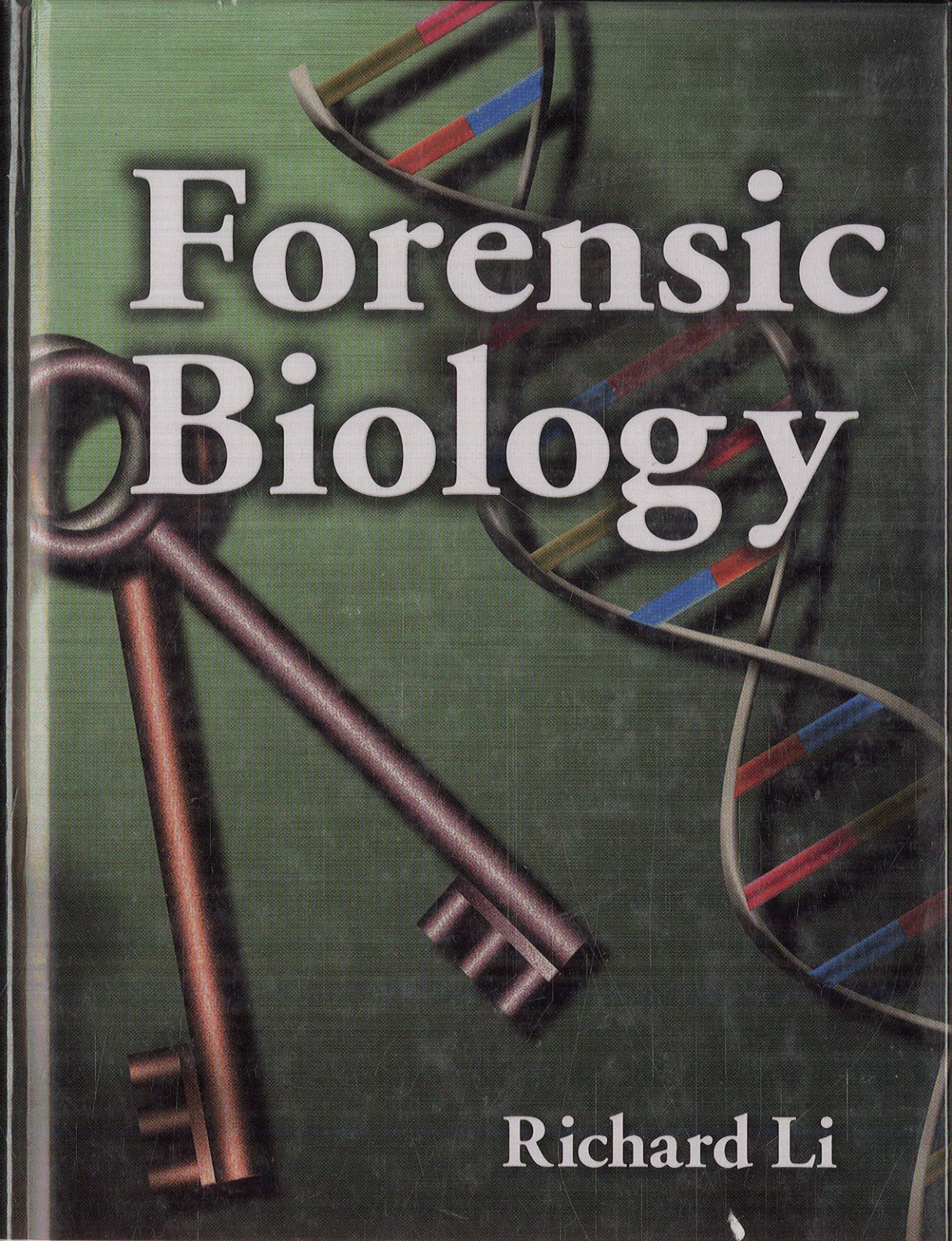


The background of the cover features a large, ornate metal key, possibly a skeleton key, positioned diagonally from the bottom left towards the center. To the right of the key, a DNA double helix structure is depicted, with its sugar-phosphate backbones in grey and its base pairs represented by red, blue, and yellow segments. The entire scene is set against a dark green, textured background.

Forensic Biology

Richard Li



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