

COMPREHENSIVE BIOCHEMISTRY

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VOLUME 19^{B/II}

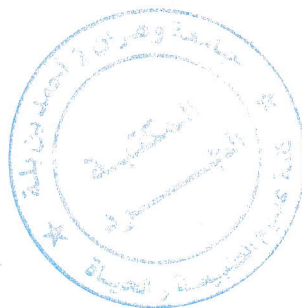
PROTEIN METABOLISM

ELSEVIER

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PROTEIN METABOLISM



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Their Structure, Function and Metabolism
 by DAVID I. STOTT and ALAN R. WILLIAMSON

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