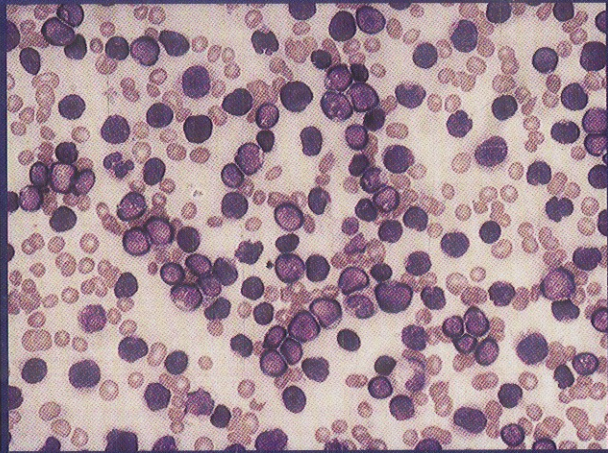
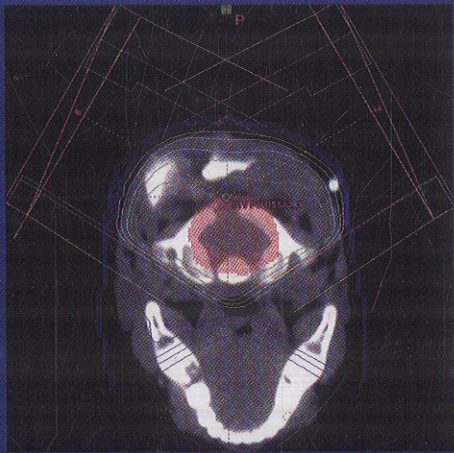


DEBORAH TOMLINSON
NANCY E. KLINE
Editors

Pediatric Oncology Nursing

PEDIATRIC ONCOLOGY



ADVANCED
CLINICAL HANDBOOK
SECOND EDITION

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