A 9 year old boy with history of cystic nephroma excised at 20 months of age (Figure 1) presented with a neck swelling. Examination revealed a 30 mm x 30 mm firm, nontender mass in the left lobe of the thyroid. There were no associated compressive or obstructive findings and no cervical lymphadenopathy. Thyroid function and thyroid autoantibody titers were normal. Ultrasound demonstrated the thyroid gland replaced by cysts and nodules (Figure 1, Supplemental Figure 1). Fine needle aspiration of the dominant nodule was categorised as indeterminate according to the Bethesda Thyroid Cytopathology Reporting System (1), however cytological features favored a colloid nodule as part of a multinodular goitre (MNG) showing papillary hyperplasia (Supplemental Figure 2).

Twelve months earlier, the proband’s father had noted a similar neck swelling and MNG was diagnosed based on clinical and radiological findings (Supplemental Figure 3). FNA demonstrated a benign cytologic pattern consistent with colloid nodules. The deceased paternal grandfather had a verbally reported but unconfirmed history of thyroidectomy. Neither the proband nor his father had thyroid surgery, electing instead to undertake regular thyroid sonographic surveillance.

Review of the proband’s chest CT scans (performed at 20 months of age), identified pulmonary cysts consistent with pleuropulmonary blastoma (PPB) type Ir. Notably, his father had been diagnosed with a lung cyst at age 13, following presentation with acute chest pain, likely also PPB type Ir (figure 2).

Both individuals provided informed consent for institutionally-approved genetic research and this case report. They were found to carry a novel germline DICER1 mutation: c.5221_5232delAACAACACCATC.

DICER1 syndrome is a recently described, highly pleiotropic, variably penetrant, autosomal dominant tumor predisposition syndrome (2), caused by inactivating germline DICER1 mutations (3). It is associated with rare cancers and dysplasias, occurring typically from birth to age 20 years: most frequently PPB (4), CN, ovarian Sertoli-Leydig cell tumors and MNG (5). Thyroid disease (nodular hyperplasia, cysts, MNG) may emerge as the most frequent presentation of this syndrome.

DICER1 is a cytoplasmic RNAse III endoribonuclease, producing mature microRNAs: small, noncoding RNAs that post-transcriptionally modulate messenger RNA (mRNA) expression and are critical in early somatic de-
velopment (6). Interestingly, DICER1 shares a locus with the familial MNG gene (MNG1) on chromosome 14q (7).

Childhood and/or familial MNG suggest DICER1 Syndrome, particularly when other highly characteristic diseases like CN or lung cysts co-occur. Genetic counseling is strongly recommended (www.ppbregistry.org) prior to DICER1 mutation analysis. The prevalence of differentiated thyroid carcinoma in DICER1-associated nodular thyroid disease is unknown, however follicular thyroid carcinoma has been reported following high dose chemotherapy and hematopoietic stem cell transplantation for treatment of PPB (8). Wider recognition of the DICER syndrome will shed light on the natural history of the associated thyroid disease and facilitate early detection of other tumors, including PPB.

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Figure 2. Computed chest tomographic images of proband (C) and his father (D) revealing pulmonary cysts consistent with PPB type Ir.

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References