Incidentally detected asplenia in a healthy 64-year-old female live kidney donor

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Abstract
Heterotaxia syndromes are rare birth defects which can result in developmental malformations. A 64-year-old woman presented to the hospital for preoperative screening for kidney donation; during which she was found to have no gallbladder and no spleen, without any signs of surgical removal. This could be a new description of a heterotaxia syndrome.

INTRODUCTION
Heterotaxia syndromes, also known as isomerism syndromes, are rare birth defects which are the result of a failure to establish normal left and right asymmetry during the development in intrauterine life [1, 2]. There are various outcomes of this congenital construction disorder, which include situs ambiguous, situs inversus, polysplenia syndrome and Ivemark syndrome [1, 3–6].

These developmental disorders can result in many malformations, such as pulmonary isomerism, four-lobed lungs, cardiac malformations, transposition of the great cardiac vessels, septal defects, midline liver, malrotation of the bowel, asplenia, renal agenesis and also an untraceable gallbladder. In short, all internal organs can be affected [3, 4]. In most of the described cases until now, anomalies in multiple organs are found. Anomalies in only two organs are a much more uncommon scenario.

CASE REPORT
A 64-year-old Dominican woman presented to the Nephrology department because she wished to donate one of her kidneys to her son, who suffered from end stage hypertensive
nephropathy. Her medical history included a laparoscopic sterilization in 1981, mammary cyst removal in 1988, a vaginal hysterectomy in 1999, a tonsillectomy in 2004 and a right-sided tympanoplasty in 2005. She had not experienced severe illnesses in the past and therefore had not received additional vaccinations. She used Tramadol and Paracetamol two to three times a week because of incidental headaches. Her family history included cardiovascular disease and diabetes mellitus.

Physical examination showed a generally healthy woman. Her body mass index was 31 and she was normotensive. Head and neck examination revealed no abnormalities. Heart and lung examination were normal. Abdominal examination revealed scars of the laparoscopic sterilization. Liver and spleen could not be palpated.

She underwent a complete workup for kidney donation. Similar to her son, she was blood group O positive. Human leucocyte antigens mismatch was 1-1-1, and the crossmatch test with her son was negative. Blood tests revealed a hypothyroidism, as well as slightly elevated HbA1c-levels. The oral glucose tolerance test was normal and her hypothyroidism was medically corrected. Urinalysis showed no abnormalities and urine culture was negative. Virology reports showed no contraindication for kidney donation. A cardiac ultrasound and exercise stress test were normal.

Finally, she had a plain chest radiography and a computed tomography scan of the abdomen. The chest X-ray showed no abnormalities. The computed tomography (CT) scan revealed no abnormalities of the kidneys. However, no gallbladder or spleen could be detected (Fig. 1) and the report mentioned a ‘post-cholecystectomy and post-splenectomy situation’. After the initial screening, she was referred to the Transplant Surgery outpatient clinic for surgical screening for live kidney donation. The discrepancy between her medical history and physical examination on one hand and the CT scan report on the other hand was noticed, but was not felt to be a contraindication for kidney donation. After multidisciplinary team discussion, she was approved as a live kidney donor. The right-sided laparoscopic donor nephrectomy was scheduled with Cefazoline prophylaxis.

During the donor nephrectomy, we explored the abdomen in order to find signs of a gallbladder or a spleen, but both organs remained untraceable (Fig. 2). Also, no accessory spleen was seen. The donor nephrectomy was successful, as was the transplant procedure in her son, and postoperative recovery was uneventful.

To check whether the patient had a real asplenia, the presence of Howell–Jolly bodies in circulating erythrocytes was investigated. Howell–Jolly bodies are nuclear remnants of red blood cells which are removed from the blood while passing the spleen. In asplenia, Howell–Jolly bodies are visible on a standard hematoxylin- and eosin-stained blood smear as purple (basophilic) spots on pink (eosinophilic) erythrocytes. The result of her test was positive, indicating a lack of splenic tissue in her body.

As no splenic tissue had been detected, and she had not received any vaccinations against pneumococci, Haemophilus Influenza-B and meningococci as she was not aware of her asplenia, she was advised to get these vaccinations and receive a yearly vaccination against the common flu.

**DISCUSSION**

In conclusion, in our live kidney donor, an absent gallbladder and asplenia was detected incidentally. The asplenia was confirmed with the presence of Howell–Jolly bodies. Although she had never suffered from severe infections with polysaccharide encapsulated bacteria, we felt that it would be wise to counsel her on vaccination and antibiotic prophylaxis, following the existing Dutch guidelines for post-splenectomy patients [7].

**Figure 1:** CT scan: no spleen can be detected in the left upper quadrant of the abdomen.

**Figure 2:** (A) View of the right upper quadrant of the abdomen during laparoscopic donor nephrectomy. Normally, the gallbladder would be found around the vertical line on the liver capsule. (B) View of the left upper quadrant of the abdomen. Part of segment 3 of the liver and part of the stomach can be seen. No spleen could be detected.
Heterotaxia syndromes have expressed themselves in various ways, but to this day no case has been described with just an absent gallbladder and asplenia. This could be a new presentation of a heterotaxia syndrome, as there are no signs of other underlying conditions.

CONFLICT OF INTEREST STATEMENT
None declared.

ETHICAL APPROVAL
Not required.

CONSENT
Yes.

GUARANTOR
Dr W.N. Nijboer.

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