The genetic aspect and morphological appearance of achondrogenesis

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Received: 21 June 2017
Accepted: 30 June 2017

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ABSTRACT

Achondrogenesis (ACG) is a number of disorders that are the most severe form of congenital chondrodysplasia characterized with bones and cartilage malformation. Generally, characteristic of ACG is a small body, short limbs, and other skeletal abnormalities. As a result of infants with ACG their serious health problems, usually born prematurely, are stillborn, or die shortly after birth from respiratory failure. Currently 3 type variants of ACG such as ACG-1A, ACG-1B and ACG-2. ACG-1A appears to be autosomal recessive (AR), with thyroid hormone receptor interactor 11 (TRIP11) gene mutation, while ACG1B also appears to be AR, with diastrophic dysplasia sulfate transporter or DTDST (SLC26A2) gene mutation. ACG-2 is caused by autosomal dominant (AD), with type 2 collagen (COL2A1) gene mutation. ACG-1A had characterized such as physis abnormal, vertebral bodies with unossified or small and oval, skull poorly ossified, periodic acid–Schiff (PAS) stain positive chondrocyte inclusions finding in long bones. ACG-1B had physis abnormal, vertebral bodies with unossified or small and oval, skull ossified, perichondrocyte collagen rings finding in long bones. ACG-2 also had physis abnormal, metaphyseal cupping, vertebral bodies with unossified or small and oval, enlarged chondrocyte lacunae.

Keywords: Achondrogenesis, COL2A1, Gen mutation Physis abnormal, SLC26A2, TRIP11

INTRODUCTION

Achondrogenesis (ACG) is a number of disorders that are the most severe form of congenital chondrodysplasia (malformation of bones and cartilage). These conditions are characterized by a small body, short limbs, and other skeletal abnormalities. As a result of their serious health problems, infants with ACG are usually born prematurely, are stillborn, or die shortly after birth from respiratory failure. Some infants, however, have lived for a while with intensive medical support.1

The other characteristic of ACG is a lethal form of congenital dystrophy characterised by micromelia macrocephaly and a short trunk that involves both proximal and distal extremity segments.2,3

Based on radiologic and histopathologic appearance, generally there are 3 variants of ACG, namely ACG type 1A (Houston-Harris type), ACG type 1B (Parenti-Fraccaro type = Fraccaro type), and ACG type 2 (Langer-Saldino type). ACG type 1A (ACG-1A) with apparently normal cartilage matrix but inclusions in chondrocytes, ACG type 1B (ACG-1B) with abnormal cartilage matrix.4 These findings were confirmed by another group shortly thereafter.5 Some cases have been published previously and diagnosed as ACG-1B, and the other cases was diagnosed as ACG-1A, while Parenti case this may be classified as ACG type 2 (ACG-2).4,6-9 ACG-1A caused by mutation of thyroid hormone receptor interactor 11 (Trip11) gene.10 Inheritance pattern of ACG-1A appears autosomal recessive (AR),10,11 ACG-1B caused by mutation of the diastrophic dysplasia sulfate transporter.
gene (SLC26A2). Inheritance pattern of ACG-1B appears AR. ACG type 2 (ACG-2) caused by mutation of the type 2 collagen gene (COL2A1). Inheritance pattern of ACG-2 appears autosomal dominant (AD).11 In 1983, a new classification of ACG was classified into 4 type and adopted in the McKusick catalogue.12,13

Not just the morphological appearance of ACG caused by genetic factors. In the other cases revealed that morphological appearance among others determined by body mass index (BMI). According to state that BMI is determined by genetic and environmental factors.14 More over, one of the genetic factors that determine the BMI is a genetic polymorphism of sex hormone binding globulin (SHBG), whereas the intake of nutrients is one of the environmental factors. In detail, that differences of dietary intake at mother in childhood age with BMI <18.5 kg/m² and has heterozygous variant SHBG genotype (W/v) determine to status of chronic energy deficiency (CED), therefore that low protein, fat and carbohydrate intake then getting low of CED status.15 We suspect that beside of genetic factor, environment factor such as dietary intake also affected to ACG. Because that too need of ACG reports from genetic aspect and dietary intake.

The genetically of achondrogenesis

ACG-1A had mutation of Trip11 gen. The Trip 11 gen encodes the Golgi microtubule-associated protein 210 (GMAP-210). The biochemical data provide strong evidence that GMAP-210 is a Golgi-associated protein that directly interacts with microtubule ends.16 The mutation of Trip 11 gen caused loss-of-function GMAP-210. GMAP-210 moves proteins from the endoplasmic reticulum to the Golgi apparatus.1 More over that mice lacking GMAP-210 die at birth with a pleiotropic phenotype that includes growth restriction, ventricular septal defects of the heart, omphalocele, and lung hypoplasia.17

The Golgi apparatus is an organelle with multiple complex functions. The Golgi complex is involved in cellular processes other than the classical trafficking and biosynthetic pathways. These organel can be considered as a cellular headquarters where cargo sorting/processing, basic metabolism, signalling and cell fate decisional processes converge.18 Because the defect of GMAP-210, thus this protein is not able to move from endoplasmic reticulum to Golgi apparatus, and they remain in the endoplasmic reticulum, which swells up. The circumstances led to disruption of skeletal formation process, so that the fetus has an abnormal phenotype. More over, the TRIP11 gene analyzed in 10 unrelated patients with ACG-1A. For all patients was done to identified of homozygous or compound heterozygous loss-of-function mutations. The result of analysis showed that two mutations (c.202-2A→G and c.589-2A→G) affect intronic splice-acceptor sites and five mutations (p.R264X, p.R1028X, p.Q1160X, p.R1167X, and p.W1224X) are nonsense mutations.1

ACG IB caused by mutations in SLC26A2 (diastrophic dysplasia sulfate transporter or DTDST) gene. The SLC26A2 gen located on 5q, hereditary by autosomal recessive (AR).13 The gen of SLC26A2 which encodes a sulfate transporter.1 More over, mutated alleles of SLC26A2 gene cause each of the four recessive chondrodysplasias, ie: diastrophic dysplasia (DTD), multiple epiphyseal dysplasia (MED), atelosteogenesis Type II (AO2), and ACG-1B. Result of study showed that SLC26A2 mRNA and protein immunostaining were detected in developing fetal hyaline cartilage. More over, SLC26A2 expression is also detected in tissues not affected in chondrodysplasias caused by SLC26A2 mutations.19

ACG-2 is characterized genetically by a mutation in the COL2A1 gene. The COL2A1 gen located on chromosome 12. Hereditary of ACG-2 by autosomal dominant (AD). The COL2A1 gene involved in the production of type 2 collagen, which is essential for hyaline cartilage formation and endochondral ossification. The mutation of COL2A1 gen affects type 2 procollagen formation, disrupting formation of the triple helix conformation necessary for proper function. Because that decreased type 2 collagen secretion and abnormal intracellular retention of the defective protein.20 The other cases showed that overmodified type 2 collagen and the presence of type 1 collagen was found in the cartilage matrix of all seven cases. Five patients were heterozygous for a nucleotide change that predicted a glycine substitution in the triple helical domain (G313S, G517V, G571A, G910C, G943S). Analysis of cartilage type 2 collagen in all five cases suggested incorporation of the abnormal α1 (II) chain in the extracellular collagen trimers. The G943S mutation has been reported previously in another unrelated patient with a strikingly similar phenotype, illustrating the possible specific effect of the mutation.21 More over also reported that histologic sections of the epiphyses show a markedly disordered physis with advanced periesteal ossification. Besides that, corresponding to radiologic metaphyseal cupping, deficient chondroid matrix, markedly enlarged chondrocyte lacunae, and increased cartilage vascularity with perivascular fibrosis.22,23

Morphological appearance of achondrogenesis type 1

The first cases of ACG-1 was detected by prenatal ultrasonography (USG) at 20 weeks gestation. The USG showed single live intrauterine with gross skeletal dysplasia-ACG-1. On USG there was absence of forearm bones in both upper limbs and absence of bones in the right lower limb. A dwarfed fetus with large head, short neck and chest and short limbs was terminated trans-vaginally. Radiologic examination revealed appearance of ACG-1. Though the case had no known risk factor and the phenotypic abnormality was mild, modern
development in prenatal screening made the early detection possible. The other data in this case showed that a detailed family history of the patient and her husband should be elicited along with clinical, genetic, radiographic and morphological examination. ACG-1 from India presented in Figure 1.


**ACG-1=A. Fetus viewed by ultrasonography. B. The radiograph of fetus. C. The photograph of fetus postmortem neonatal.**

**Figure 1. Achondrogenesis type 1 from India.**

The second cases of ACG-1 reported that 32-year-old Turkish woman, gravida 3, para 2. Her patient diagnoses of gestational diabetes mellitus, polyhydramnios, and abnormal results of an ultrasound examination performed by obstetrician. First degree consanguinity was noted between the couple. Her first pregnancy had ended in a cesarean section at 33 weeks' gestation with a diagnosis of severe preeclampsia and breech presentation. Although it had not been possible to make an exact diagnosis, the appearance of the male baby had suggested dwarfism.

![Table 1: Prenatal sonograms examination and characteristic of the fetus achondrogenesis type 1 from Turkish women.]

**Table 1: Prenatal sonograms examination and characteristic of the fetus achondrogenesis type 1 from Turkish women.**

<table>
<thead>
<tr>
<th>Prenatal sonograms examination</th>
<th>Characteristis of the fetus was terminated by cesarean section at 34 weeks and 4 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Micromelia, narrow thorax, high abdomen/thorax ratio, pulmonary hypoplasia, and poor mineralization of the skull and vertebrae.</td>
<td>- Female baby died within the first thirty minutes of birth.</td>
</tr>
<tr>
<td>- Polyhydramnios and a pseudohydric appearance, pleural and pericardial effusion, and low-lying ears</td>
<td>- Birth body weight 1810 gram and 31 centimeters.</td>
</tr>
<tr>
<td></td>
<td>- Hydrops, severe tetramicromelia, and a flat face.</td>
</tr>
<tr>
<td></td>
<td>- The head was disproportionately large relative to the reduced neck, trunk and limb length.</td>
</tr>
<tr>
<td></td>
<td>- Anteroposterior and lateral postmortem whole-body radiographs revealed inadequate ossification of the bones, except for the clavicles, and there were short beaded ribs with multi-ple fractures, minimal ossification of vertebral bodies, arched iliac wings, stellate femurs and humeri, and micromelic long bones.</td>
</tr>
</tbody>
</table>

**Achondrogenesis type 1A**

ACG-1A derived from Turkey, pregnant women gravida 2, parity 1, consult regularly to Obstetricians. The examination was conducted using ultrasonography at week 16 of the last menstrual period. The examination results indicate that the presence of skeletal dysplasia. No characteristics in a personal or family history. Laboratory results were in the normal range. Using the ultrasonography shows a single living fetus consistent with the 17th gestational week in terms of head circumference, biparietal diameter and abdomen circumference and with the 12th gestational week in terms of femur length. During the examination by USG, the case was established to have distinct retardation in all long bones when compared to those of the same
gestational age in addition to macrocephaly, narrow thorax, protuberant abdomen, reduced thorax/abdomen ratio, reduced vertebral echogenicities, horizontally placed short ribs with irregular cortex and increased echogenicity, pyelectasis, protruding forehead, nasal bone hypoplasia, increased skin thickness, and polyhydramnios.

A. The radiograph of the fetus. The fetus shows shortness of extremities and reduced ossification in calvarium and vertebrae. B. Post-termination photograph of the fetus. The long bones of all extremities appear to be short, and macrocephaly, flat nasal root, protruding forehead, low-set ears and narrow thorax are prominent.

**Figure 3: The photograph and radiograph of achondrogenesis type 1A from Turkey.**

The result of amniosynthesis shows that karyotype 46 XX and normal. After genetic counselling, then it is advisable for the termination of the pregnancy. Pregnancy was terminated after obtaining the consent given by the family. Post-mortem examination revealed severe micromelia, narrow thorax, abdomen wide, protruding forehead, flat nasal root, hypertelorism, and low-set ears in the fetus. They showed almost non-existent ossification in the vertebræ and loss of mineralization in the calvarium, as well as distinct shortness of the extremities. Based on the results of clinical and radiological evaluations, the fetus was diagnosed as ACG-1A. The photograph and radiograph results of ACG-1A from Turkey presented in Figure 3.

The other cases of ACG-1A was reported that human fetus at 27-weeks-old with reveals a lack of mineralization in the skull and the vertebral column and short limbs (Figure 4). The other cases reported that patient of ACG-1A showed swollen endoplasmic reticulum and vesicular Golgi apparatus. This appearance corresponds to the absence of vertebral-body and skull ossification on radiography, the lack of organized columnar zones of proliferating chondrocytes on histologic analysis. The reduced expression of Col10a1 on immunohistochemical analysis, and the expanded endoplasmic reticulum cisternae in chondrocytes on electron microscopy have all been reported in affected patients.

A. The radiograph of human fetus with reveals a lack of mineralization in the skull and the vertebral column (red arrows) and short limbs. B. The electron micrographs show chondrocytes from two unrelated fetuses with ACG-1A, with swollen endoplasmic reticulum (red arrows) and vesicular Golgi apparatus (arrowhead in upper image).

**Figure 4: Achondrogenesis type 1A of fetus showed at 27-weeks-old.**

**Achondrogenesis type 1B**

Superti-Furga reported the cases of ACG-1B with characterized in a breech position. The fetus after birth is regarded as abnormal. Fetuses show disproportion between the head, seen approaching a normal or normal size, and the rest of the body, so looked short of normal. The other appearance of fetus ie: flat face, short neck, and the neck has a soft tissue may be thickened, narrow thorax with protuberant abdomen. The fetus newborns appear fat or hydroptic, this is due to the amount of soft tissue relative to the short skeleton. Often suffer of hernia inguinal or umbilical.

**Figure 5: The result of X ray and clinical appearance of fetus with achondrogenesis type 1B.** A. The radiograph of fetus. B. Clinical appearance of fetus.
Display short femur seen during an ultrasound during pregnancy. The use of USG also visible nuchal edema, reduce the length of the buttocks, poor ossification of the vertebral body and leg bones (leading to difficulty in determining reviews in length), and polyhydramnios. Fetus with ACG-1B born at 34 weeks and died 25 minutes after birth.13 The view of X ray and clinical appearance of ACG-1B presented at Figure 5.

Clinical appearance of achondrogenesis type 2

Female human fetus was delivered transvaginally at 22 weeks gestation and died shortly after birth. Until 20 weeks of gestation showed uneventful, when an USG examination showed that the fetus significantly shortened femur for the gestational age and nuchal edema.

Table 2: The characteristic of morphological appearance, radiological and histological examination of fetus was diagnosis achondrogenesis type 2.

<table>
<thead>
<tr>
<th>Morphological appearance</th>
<th>Radiological</th>
<th>Histological</th>
</tr>
</thead>
<tbody>
<tr>
<td>-Fetus body-weight: 490 grams.</td>
<td>-Shortness of limbs. The tubular bones were short and broad. The metaphyses of tubular bones were widened with irregular ends and lateral spurs.</td>
<td>-Chondroosseous tissues showed hypervascularity of cartilage. Moreover, cellular density was high and the matrix was reduced.</td>
</tr>
<tr>
<td>-Body length of 17.3 cm</td>
<td>-The calvaria were relatively well ossified.</td>
<td>-The cells were large, often starred, and their lacunae were enlarged.</td>
</tr>
<tr>
<td>-She had a craniofacial anomaly ie relatively large calvarium. The size of the head circumference ie 22.3 cm and biparietal diameter ie 06.15 cm.</td>
<td>-The vertebral bodies were insufficiently ossified, predominantly in the cervical spine.</td>
<td>-In the growth zone, the cells showed irregular columns. Moreover, this cell ended in hypertrophic cells placed irregularly.</td>
</tr>
<tr>
<td>-Micrognathia, small mouth, tongue hypertrophy, flat nose and hypertelorism.</td>
<td>-The ribs were horizontal without fracture.</td>
<td>-Vascular penetration was irregular in the ossification line. The primary trabeculae were thickened, irregular and uneven.</td>
</tr>
<tr>
<td>-The neck of the fetus is very short, so short chest with antero-posterior flattening.</td>
<td>-The iliac wings were square with a horizontal acetabular angle and a concave internal edge.</td>
<td></td>
</tr>
<tr>
<td>-The abdomen was large and distended. There was severe shortness of upper and lower limbs, especially from rhibzoemetic segment.</td>
<td>-The ischiatic and pubic bones were nonossified.</td>
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<tr>
<td>-Extremities are also bent and bilateral club foot.</td>
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</tbody>
</table>

The fetus was conceived by a 26-year-old mother, while her husband was 30 years old. In the family can not be found ACG history and are not known exposure of the teratogens. The results of amniocentesis showed that the baby is female and has a normal karyotype.

Based on examination of organs revealed that the growth and development of gestational age there are no significant anomalies. On the basis of clinical findings, radiological and microscopic, this case was diagnosed as a mild form of ACG-2 (Langer-Saldino).27 The characteristic of morphological appearance, radiologic and histologic examination of fetus was diagnosis ACG-2 presented in Table 2.

The examination by ultrasonography, radiography, photography and microscopy of the fetus of ACG-2 presented in Figure 6.

Figure 6: Examination by ultrasonography, radiography, photography and microscopy of the fetus of achondrogenesis type 2.
**Achondrogenesis type 2 at the couple had a fourth pregnancy**

A familial case of ACG-2 caused by COL2A1 gene mutation and ‘patchy’ expression in the mosaic father was reported. The case of ACG-2 occurred on nonconsanguineous young couple had four pregnancies of an apparently healthy. The first child was born at 32 weeks and died neonatally. In the second pregnancy at 17 weeks gestation showed that short limbs and fetal hygroma in USG examination. Similar findings were observed in the third fetus. These couple had a fourth pregnancy showed findings consistent with ACG-2.28

The characteristic of ACG-2 at the couple had a fourth pregnancy presented at Table 3. The result of examination by USG for fourth fetus with ACG-2 at the couple had a fourth pregnancy presented at Figure 7. The other cases of ACG-2-hypochondrogenesis and severe spondyloepiphyseal dysplasia congenita (SED) are lethal forms of dwarfism caused by dominant mutations in type 2 collagen gen (COL2A1). In these cases, reported that 7 patients had mutation of COL2A1 gen. For all 7 patients, 2 patients were male and 5 females. This happens because that ACG-2 are autosomal dominant. The seven patient had body length varies ranging from 29 cm to 39 cm on gestational age between 35-42 weeks, short limbs dan small thorax. Two of 7 patients had cleft palate, 6 of 7 patients had respiratory insufficiency at birth. Age of death varies from 6 hours to 1 year 1 month.

**Figure 7: The result of examination by USG for fourth fetus with achondrogenesis type 2 at the couple had a fourth pregnancy.**

**Table 3: The characteristic of achondrogenesis type 2 at the couple had a fourth pregnancy.**

<table>
<thead>
<tr>
<th>The first pregnancy</th>
<th>The second pregnancy</th>
<th>The third pregnancy</th>
<th>The fourth pregnancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal age 30 years.</td>
<td>Maternal age 32 years.</td>
<td>Maternal age 33 years.</td>
<td>Screened at 10.6 weeks for prenatal diagnosis by chorionic villus sampling.</td>
</tr>
<tr>
<td>Evaluated at 28 weeks and 5 days for suspected abnormal development of fetal limbs and mild polyhydramnios.</td>
<td>Evaluated at 16 weeks of pregnancy for fetal karyotyping as 46, XX. At USG, the fetus showed a large and septated nuchal cystic hygroma associated with hydrops and hydropsic bowel.</td>
<td>Evaluated for fetal karyotyping at 17+3/7 weeks.</td>
<td>Transvaginal USG showed a septated nuchal cystic hygroma with enlarged jugular lymphatic sacs, generalized fetal hydrops developed 1 week later.</td>
</tr>
<tr>
<td>Examination at 28 weeks with USG showed a viable fetus was recognized with circumferences of the head and abdomen appropriate for gestational age. All long bones showed that the limb shortening involved all segments. The other anomalies as: micrognathia, fetal skin redundancy in the neck and thorax region, and a ‘bell-shaped’ thorax. The spine had an ovoid shape and flattening of vertebral bodies.</td>
<td>Fetal biometry of the head was appropriate for gestational age.</td>
<td>The circumference of the head and abdomen were appropriate for gestational age.</td>
<td>Amniotic fluid, fetal biometry and heart rate were normal.</td>
</tr>
<tr>
<td>The pregnancy ended with preterm labor at 32-week gestation. Weight of fetus a 1.700 gram, male and died in the newborn period. There was a short, thick neck, flat face with a deep nasal bridge, micrognathia, and short limbs.</td>
<td>Micrognathia.</td>
<td>All long bones showed that only the proximal segments were shortened.</td>
<td>The pregnancy was terminated at 18 weeks gestation.</td>
</tr>
</tbody>
</table>

Abbreviations: USG = ultrasonography.
Resting cartilage with mild to very hypervascular and slightly increased vascular canals. Chondrocytes with inclusion bodies and 4 of 7 patients showed dilated rough endoplasmic reticulum. Six of 7 patients with abnormal growth palate. The radiographs of patients ACG-2 and severe spondyloepiphyseal dysplasia congenita presented in Figure 8.

A. Patient R86-153 represents a severe phenotype characterised by short tubular bones with metaphyseal irregularities, minimal ossification of vertebral bodies in the thoracic spine, hypoplastic iliac wings with flat acetabular roofs, and short ribs. B. Patient R83-32 shows an intermediate phenotype. C. Patient R91-68 illustrates a milder phenotype with normal metaphysis of shortened tubular bones, flattened but normally ossified vertebral bodies, normal acetabular roofs, and a less narrow chest.

**Figure 8: The radiographs of patients with achondrogenesis type 2 and severe spondyloepiphyseal dysplasia congenita.**

**Achondrogenesis type 2 in a fragmented fetus**

The cases of ACG-2 in a fragmented fetus was reported. These cases experienced by mother of age 29 years, gravida 2 para 1 with a history of gestational diabetes, asthma, smoking, and prior cesarean section.

A. The radiographs of the upper extremities show markedly shortened limbs and long bones. The metaphyseal ends, especially of the distal humerus, show prominent metaphyseal cupping, a feature seen in ACG-2. B. The histology of the long bone metaphyses shows disorganized endochondral ossification and ballooning chondrocytes.

**Figure 9: The radiographs and histology of the long bone in achondrogenesis type 2.**

There was no relevant family history. Elective termination by dilation and evacuation (D and E) was performed at a 15 4/7 weeks. Prenatal, post mortem, radiography and histology examination of fragmented fetus in ACG-2 presented in Table 4. The radiographs and histology of the long bone in ACG-2 presented in Figure 9. The photograph, radiograph and histology examination of vertebral column in ACG-2 presented in Figure 10.

A. The photograph of the fragmented vertebral column. B. The tissue radiograph of the fragmented vertebral column. In this figure showed that the lack of ossification of the vertebral bodies and no rib fractures. C, D. Histology examination to confirms of the lack of vertebral body ossification.

**Figure 10: The photograph, radiograph and histology examination of vertebral column in achondrogenesis type 2.**

According to Hansen et al, the size toe–heel length 1.4 cm is small for 13-14 weeks of pregnancy. Based on the expected value by Elejalde and de Elejalde that the length of femora 12.5 to 27 mm, and humeri 12.5 to 26 mm, thus the length of femora 6 mm, whereas the humeri 7 mm in fragmented fetus was shorten.

Based on the findings of morphological observation, radiological, histological and genetic analysis, the fetus is fragmented concluded experiencing ACG-2. Moreover, revealed that the fetus showed heterozygous mutation in the COL2A1 gene [c.1358G>A transition (Gly453Asp) in exon 19.

Superti-Furga et al to state that a distinction between ACG-1A, 1B ACG, and ACG-2 on clinical grounds is difficult.

More details that almost normal hands are seen in ACG-2, whereas in ACG-1A and ACG-1B showed that the hands are evidently shortened. While the radiological observation showed that rib fractures, thus may suggest as ACG-1A. Guidelines for the diagnostic of ACG presented in Table 5.
Table 4: Prenatal, post mortem, radiography and histology examination of fragmented fetus in achondrogenesis type 2.

<table>
<thead>
<tr>
<th>Prenatal examination</th>
<th>Post mortem examination</th>
<th>Photograph and radiography examination</th>
<th>Histology examination</th>
</tr>
</thead>
<tbody>
<tr>
<td>USG revealed a cystic hygroma, possible facial clefting, an abnormal 4-chamber heart, a small chest, upper limb shortening, and an unspecified lower limb deformity.</td>
<td>-Toe–heel length 1.4 cm. -Disruption seen on the head and skull bones, thus preventing the determination of the head circumference. The right eye seemed distracted, as well as the left ear appeared posteriorly rotated. -Disrupted showed to the spinal column, thoracic cavity, diaphragm, pericardial cavity and abdominal cavity, and some of the organs were not identified among them any loose fragments. -Partially disrupted also showed on the liver, pancreas, heart, and intestines. The placenta was fragmented. -Disrupted partially also showed on extremities, more over for all 4 extremities were markedly shortened. The right hand was intact and showed mild clinodactyly of the 5th finger but no polydactyly or syndactyly. On the other hand, the left hand was partially disrupted. -The internal genitalia were disrupted, and the external genitalia were female. The anus could not be identified.</td>
<td>- The remains fetus showed ossification, but it is clear that the long bones shorten with no fractures. - The length of femora 6 mm, whereas the humeri 7 mm, with metaphyseal cupping of both femora and humeri was seen. - No ossification of the vertebral body showed by photograph and radiograph.</td>
<td>- Disordered physis with advanced periosteal ossification, chondroid deficient matrix, enlarged chondrocyte lacunae, and increased cartilage vascularity with perivascular fibrosis. - The lack of vertebral body ossification, but skull ossification was present and also rib ossification was present without fractures. - No chondrocyte inclusions. - Disrupted markedly on the pelvic bones, while the hands showed faint ossification of all phalanges. The distal femoral epiphysis and proximal tibial epiphysis were not ossified.</td>
</tr>
</tbody>
</table>

USG = ultrasonography.

Table 5. Guidelines for the diagnostic of achondrogenesis.

<table>
<thead>
<tr>
<th>Part of skeleton</th>
<th>Extremely short femora and humeri with</th>
<th>Additional skeleton features</th>
<th>Suggested diagnosis</th>
<th>Key histologic findings in long bones</th>
<th>Mode of inheritance</th>
<th>Genetic mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Long bones</td>
<td>Pattern: Metaphyseal cupping also look for: unossified, small or oval vertebral bodies</td>
<td>-</td>
<td>ACG-2</td>
<td>- Physis abnormal -Markedly enlarged chondrocyte lacunae</td>
<td>Autosomal dominant</td>
<td>COL2A1</td>
</tr>
<tr>
<td></td>
<td>Pattern: None of the above specific long bone findings</td>
<td>Vertebral bodies: unossified or small and oval</td>
<td>Skull poorly ossified</td>
<td>ACG-1A</td>
<td>- Physis abnormal - PAS + chondrocyte inclusions</td>
<td>Autosomal recessive</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Skull ossified</td>
<td>ACG-1B</td>
<td>- Physis abnormal - Perichondrocyte collagen rings</td>
<td></td>
</tr>
</tbody>
</table>

CONCLUSION

Base on the fact of many case report we concluded that gen mutation caused abnormality in protein and cell metabolism. Abnormality in protein and cell metabolism than caused many organs disorders form, characterized with bones and cartilage malformation, and finally caused of congenital chondrodysplasia. Currently 3 type of ACG, such as ACG-1A, ACG-1B and ACG-2. ACG-1A caused by TRIP11 gen mutation, while ACG1B caused by diastrophic dysplasia sulfate transporter or DTDS (SLC26A2) gen mutation, whereas ACG-2 caused type 2 collagen (COL2A1) gen mutation. ACG-1A had characterized such as physis abnormal, vertebral bodies with unossified or small and oval, skull poorly ossified, chondrocyte inclusions finding in long bones. ACG-1B had physis abnormal, vertebral bodies with unossified or small and oval, skull ossified, perichondrocyte collagen...
rings finding in long bones. ACG-2 also had physis abnormal, metaphyseal cupping, vertebral bodies with unossified or small and oval, enlarged chondrocyte lacunae.

**Funding:** No funding sources  
**Conflict of interest:** None declared  
**Ethical approval:** Not required

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Cite this article as: Parwanto MLE. The genetic aspect and morphological appearance of achondrogenesis. Int J Reprod Contracept Obstet Gynecol 2017;6:3203-12.