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Product datasheet for TA330536

PITX2 Rabbit Polyclonal Antibody

Product data:

| Product Type: | Primary Antibodies |
|-------------------------|--|
| Applications: | WB |
| Recommended Dilution: | WB |
| Reactivity: | Human |
| Host: | Rabbit |
| lsotype: | IgG |
| Clonality: | Polyclonal |
| Immunogen: | The immunogen for anti-PITX2 antibody: synthetic peptide directed towards the N terminal of human PITX2. Synthetic peptide located within the following region: METNCRKLVSACVQLGVQPAAVECLFSKDSEIKKVEFTDSPESRKEAASS |
| Formulation: | Liquid. Purified antibody supplied in 1x PBS buffer with 0.09% (w/v) sodium azide and 2% sucrose. Note that this product is shipped as lyophilized powder to China customers. |
| Conjugation: | Unconjugated |
| Storage: | Store at -20°C as received. |
| Stability: | Stable for 12 months from date of receipt. |
| Predicted Protein Size: | 35 kDa |
| Gene Name: | paired like homeodomain 2 |
| Database Link: | <u>NP_700475</u> <u>Entrez Gene 5308 Human</u> <u>Q99697</u> |



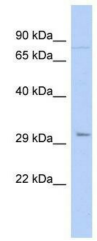
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PITX2 Rabbit Polyclonal Antibody – TA330536

| Background: | The PITX2 gene encodes a member of the RIEG/PITX homeobox family, which is in the bicoid class of homeodomain proteins. This protein acts as a transcription factor and regulates procollagen lysyl hydroxylase gene expression. Mutations in PITX2 are associated with Axenfeld-Rieger syndrome (ARS), iridogoniodysgenesis syndrome (IGDS), and sporadic cases of Peters anomaly. This protein is involved in the development of the eye, tooth and abdominal organs. It also acts as a transcriptional regulator involved in basal and hormone-regulated activity of prolactin. This gene encodes a member of the RIEG/PITX homeobox family, which is in the bicoid class of homeodomain proteins. The encoded protein acts as a transcription factor and regulates procollagen lysyl hydroxylase gene expression. This protein plays a role in the terminal differentiation of somatotroph and lactotroph cell phenotypes, is involved in the development of the eye, tooth and abdominal regulator involved in basal and hormone-regulated activity of prolactin. Mutations in this gene are associated with Axenfeld-Rieger syndrome, iridogoniodysgenesis syndrome, and sporadic cases of Peters anomaly. A similar protein in other vertebrates is involved in the determination of left-right asymmetry during development. Alternatively spliced transcript variants encoding distinct isoforms have been described. |
|-------------|--|
| Synonyms: | ARP1; Brx1; IDG2; IGDS; IGDS2; IHG2; IRID2; Otlx2; PTX2; RGS; RIEG; RIEG1; RS |
| Note: | Horse: 100%; Human: 100%; Mouse: 93%; Pig: 92%; Guinea pig: 92% |

- Protein Families: Transcription Factors
- Protein Pathways: TGF-beta signaling pathway

Product images:



WB Suggested Anti-PITX2 Antibody Titration: 0.2-1 ug/ml; ELISA Titer: 1:62500; Positive Control: COLO205 cell lysatePITX2 is supported by BioGPS gene expression data to be expressed in COLO205

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