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Published in:
Zoological research

DOI:
10.24272/j.issn.2095-8137.2022.086

Publication date:
2022

Document version
Publisher's PDF, also known as Version of record

Document license:
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Citation for published version (APA):
Single-cell transcriptomic landscape of the sheep rumen provides insights into physiological programming development and adaptation of digestive strategies

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ABSTRACT

As an important evolutionary innovation and unique organ, the rumen has played a crucial role in ruminant adaptation to complex ecological environments. However, the cellular basis of its complex morphology and function remains largely unknown. In this study, we identified eight major cell types from seven representative prenatal and postnatal rumen samples using ~56 600 single-cell transcriptomes. We captured the dynamic changes and high heterogeneity in cellular and molecular profiles before, during, and after the appearance of keratinization and identified TBX3 as a potential marker gene. Importantly, enriched stratum spinosum cells played crucial roles in volatile fatty acid metabolism.

Received: 27 May 2022; Accepted: 27 June 2022; Online: 04 July 2022

Foundation items: This work was supported by the Strategic Priority Research Program of the Chinese Academy of Sciences (XDA26040301-02), National Natural Science Foundation of China (31970392, 32172752, 32122083), Research Funds for Interdisciplinary Subject of NWPU (19SH030408), Funds Awarded by the 1000 Talent Project of Shaanxi Province, Project for Top Young Talents Program of College of Animal Science and Technology of Nanjing Agricultural University (DKQB201904), and Talents Team Construction Fund of Jilin Agricultural University.

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acid (VFA) metabolism and immune response. Our results provide a comprehensive transcriptional landscape of rumen development at single-cell resolution, as well as valuable insight into the interactions between dietary metabolism and the rumen.

**Keywords:** Rumen; scRNA-seq; Ruminal epithelium; Keratinization

### INTRODUCTION

Ruminants are among the most successful mammalian lineages in animal evolutionary history (Chen et al., 2019). They possess remarkable and distinct multi-chambered stomachs, comprised of the rumen, reticulum, omasum, and abomasum, which collectively form a highly sophisticated natural bioreactor (Lei et al., 2018). This arrangement allows ruminants to utilize plant fiber more efficiently than other mammals, which may have led to their increasing success as extensive swathes of forest were gradually replaced by grassland around 40–50 million years ago (Chen et al., 2019).

Consumed plant material is degraded by the dense and diverse microbiota in the rumen into precursors of essential metabolites and metabolic cofactors (particularly volatile fatty acids (VFAs), microbial proteins, and vitamins) used for the production of high-value meat and milk (Xu et al., 2021). Thus, the rumen and associated microbiota play key roles not only in ruminant growth, survival, and reproduction, but also in the provision of highly nutritious food for many human societies (Mizrahi & Jami, 2018). In this symbiotic system, rumen tissue has evolved a series of distinct anatomical features and innovative functions that enable adaptation to the vast, complex rumen microbial ecosystem (Fonty et al., 1991; Russell & Rychlik, 2001). Therefore, elucidating the cellular biological mechanisms underlying rumen formation and development is crucial for understanding the origin and evolution of the organ.

Unlike the stomachs of humans and monogastric animals (with simple, glandular, and prismatic epithelium), rumen tissue has a complex structure consisting of stratified squamous epithelium, lamina propria, mucousa, tunica muscularis, and plasma membrane (Groenewald, 1993; Steven et al., 1970). In contrast to the smooth, non-keratinized stratified squamous epithelium of the esophagus (Oshima et al., 2011), the stratified squamous epithelium of the rumen, including the stratum corneum, granulosum, spinosum, and basal layer, is keratinized like skin epidermis (Graham & Simmons, 2005). This notable structural feature of the rumen not only plays a protective role in defense against microbial invasion but also facilitates renewal of rumen tissue (Graham & Simmons, 2005). This suggests that immunoregulation in the rumen may be involved in maintaining homeostasis, although the idea of the rumen as an immune-related organ remains controversial. Previous transcriptomic analyses have suggested that the KRT4 gene and keratinocyte differentiation pathway play key roles in stratum corneum formation (Bragulia & Homberger, 2009; Pan et al., 2020). The stratified squamous epithelium is the main site of VFA metabolism and ketogenesis in adult ruminants, thereby affecting the availability of energy and nutrients for the host (Rémond et al., 1995). Immunohistochemical and morphometric analyses have indicated that Na^+-K^-ATPase α-subunit expression and mitochondrial density and distribution in the stratum spinosum and basal layer of the rumen epithelium facilitate VFA metabolism (Giesecke et al., 1979; Graham & Simmons, 2005). In addition, the rumen contains diverse and interdependent populations of bacteria, archaea, protozoa, fungi, and viruses (Russell & Rychlik, 2001). However, the cellular basis of its complex morphological structure and innovative functions remains largely unknown.

Studying the development of an organ is an effective way to understand its cellular biological foundation (Finnegan et al., 2019; Haber et al., 2017). Morphometric analysis has shown that embryonic sheep rumen tissue forms three differentiated layers at ~42 days of gestation, with the papillary structure and keratinized stratified squamous epithelium forming at ~57 days of gestation (Franco et al., 2011). Postnatal rumen development differs from that of the stomach in monogastric animals. In neonatal ruminants, the fluid (milk or milk replacer) obtained from suckling primarily flows into the abomasum via the esophageal groove in the rumen, resulting in a lack of rumen microbial fermentation substrates to stimulate rumen development (Beharka et al., 1998; Wise & Anderson, 1939). After the introduction of solid food, rumen papillae continue to grow and the VFA metabolism machinery gradually matures (Baldwin & Connor, 2017; Pazoki et al., 2017; Rémond et al., 1995). These findings suggest that dramatic structural and functional changes occur during embryonic and postnatal rumen development, implying that cell type and composition have significantly changed. However, the critical cellular processes that occur during these developmental periods remain unclear.

Here, we performed single-cell RNA sequencing (scRNA-seq) to explore the cellular heterogeneity and function of specific cell types in rumen development. We profiled ~56,600 single-cell transcriptomes from seven key points in sheep rumen development (spanning prenatal to postnatal stages) to address the following questions. First, which cells form the complex morphological structure of the rumen? Second, what processes are involved in the formation of each cell type? Third, which cells play fundamental roles in specific functions and features (keratinization, papilla formation, VFA metabolism, and immunological functions) of the rumen? Our findings should help elucidate the composition, development, and functional maintenance of the rumen at the cellular level and provide valuable insight into the interactions between dietary metabolism and rumen function.

### MATERIALS AND METHODS

**Sheep rumen dissection**

The experimental design and procedures were approved by the Animal Care and Use Committee of Nanjing Agricultural University. Domesticated Hu sheep (a native breed in China with high reproductive efficiency and the ability to tolerate large amounts of roughage) were selected. Prenatal rumen samples were collected from healthy pregnant Hu sheep (at...
30, 60, 90, 110, and 130 days of gestation) via caesarean section, and postnatal rumen samples were obtained from newborn and 45-day-old lambs. Fetuses and lambs were stunned by electric shock and killed by exsanguination. The whole rumen was immediately excised and placed in ice-cold phosphate-buffered saline (PBS) (Sigma-Aldrich, USA), after which fatty tissue and blood vessels on the outer wall were carefully removed.

Primary cell isolation
Each rumen sample was incised along the dorsal curvature and washed in cold Hanks’ Balanced Salt Solution (HBSS, Gibco, USA) several times to remove mucus and blood. After collecting samples for morphological observation, the rumen tissue was cut into small pieces (5 mm), washed with HBSS until clear, and repeatedly digested with 0.25% trypsin (Gibco, USA) for 25 min at 37 °C. The digestion solution was changed every 5 min, collected in a centrifuge tube containing fetal calf serum, and filtered through a cell strainer (40 μm, Gibco, USA). After solution centrifugation (3 min at 1 500 r/min, 4°C), the sediment was resuspended in PBS to remove residual trypsin. The fetal rumen cells were centrifuged (3 min at 1 500 r/min, 4°C) again and resuspended in PBS to determine density and viability using a Bio-Rad TC20 Automated Cell Counter (Bio-Rad, USA). Cell viability was confirmed by trypan blue staining, and cell density was adjusted to 1 000 000 cells/mL. Finally, live cells were selected for scRNA-seq using the 10x Genomics single-cell platform.

scRNA-seq data processing
Single-cell raw data were processed using 10x Cell Ranger software (v3.0), including “cellranger mkfastq” to demultiplex raw data into Fastq files and “cellranger count” for alignment, filtering, barcode counting, and unique molecular identifier (UMI) counting using default settings. A sheep reference genome (Oar_rambouillet_v1.0) was downloaded from ENSEMBL and built with “cellranger mkgtf”. Finally, we obtained seven UMI count tables (absolute numbers of observed transcripts) for further analysis, i.e., one for each developmental stage.

Analysis and visualization of processed sequencing data
For downstream analyses, we retained cells expressing at least 200 genes but less than 25 mitochondrial genes. To obtain a single-cell expression matrix for the retained cells, we applied the “FindIntegrationAnchors” and “IntegrateData” functions in Seurat (v3.1.4) to merge the data and used the “ScaleData” function to reorganize mitochondrial expression. We then used the “RunPCA” function in Seurat for principal component analysis (PCA) of the corrected cell expression matrix. The number of significant principal components (PCs) was determined using the permutationPA function in the jackstraw R package (https://cran.r-project.org/web/packages/jackstraw). Analysis identified 45 significant PCs, the scores of which were used for further analysis. We then applied the “FindClusters” function in Seurat for cell clustering analysis. We set the parameter resolution to 1.2 and identified 31 clusters. Uniform Manifold Approximation and Projection (UMAP) and t-distributed Stochastic Neighbor Embedding (t-SNE) plots of the second-level clustering results were obtained with Seurat, as above.

Identification of cell types and differentially expressed genes (DEGs)
Differences in gene expression among cell clusters were analysed using the “FindMarkers” function in Seurat. The marker genes visualized in Figure 1C were identified using the FindAllMarkers Seurat function with default settings. The marker genes were analyzed using David (https://david.ncifcrf.gov) for Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment analyses.

Single-cell pseudo-time trajectory analysis
Monocle (v2.14.0) was used for ordering single cells in pseudo-time (http://cole-trapnell-lab.github.io/monocle-release/docs/#constructing-single-cell-trajectories). We constructed single-cell differentiation trajectories using Monocle-identified variable genes as ordering genes. Root state was determined based on Seurat cell identity information. Gene clusters were further divided into four clusters according to their k-means. To investigate gene function for each gene cluster, we performed GO and KEGG enrichment analyses using David (https://david.ncifcrf.gov).

Statistical analysis
Statistical analyses were performed using the R package. P-values were calculated using two-tailed Student’s t-tests, with values below 0.05 considered statistically significant.

RESULTS AND DISCUSSION
Identification of diverse cell types in rumen
To investigate the key features of rumen development, we collected seven representative Hu sheep rumen samples from the prenatal stage to functional maturity: i.e., at 30, 60, 90, 110, and 130 days of gestation, newborn (at 144 days of gestation), and 45 days old (Figure 1A). We profiled ~56 600 single-cell transcriptomes from these samples and partitioned the cells into 31 clusters (Figure 1B), each containing 194 to 4 291 cells. Based on the distinct expression patterns and levels of known cell-type marker genes (Figure 1C; Supplementary Table S1), we classified these clusters into eight cell types, including epithelial, fibroblast, smooth muscle, endothelial, proliferating, lymphocyte, macrophage, and T cells (Figure 1D).

Epithelial cells, the most variable cell population during rumen development, including basal cells, keratinocytes, differentiating keratinocytes (DKs), terminally differentiated keratinocytes (TKDs), and special spinous cells (SSCs), rapidly increased in proportion at 60 days of gestation (Figure 1E). Clusters 0, 7, and 9 were classified as basal cells with high expression of marker genes such as KRT15 and COL17A1 (Joost et al., 2016) (Figure 1E). Clusters 0, 7, and 9 were classified as basal cells with high expression of marker genes such as KRT15 and COL17A1 (Joost et al., 2016) (Figure 1E). Clusters 0, 7, and 9 were classified as basal cells with high expression of marker genes such as KRT15 and COL17A1 (Joost et al., 2016) (Figure 1E). Clusters 0, 7, and 9 were classified as basal cells with high expression of marker genes such as KRT15 and COL17A1 (Joost et al., 2016) (Figure 1E). Clusters 0, 7, and 9 were classified as basal cells with high expression of marker genes such as KRT15 and COL17A1 (Joost et al., 2016) (Figure 1E).
Ogawa et al., 2011). Clusters 14, 15, and 16 were classified as TDKs defined by high expression of CNFN and KRT23 (Figure 1C), which are associated with the cornified cell envelope of the stratified squamous epithelium (Wagner et al., 2019). Clusters 1, 3, 5, 6, and 13 were identified as SSCs with high expression of lipid metabolism and ion transport-related genes CA1, ACA2, and BDH1 (Bond et al., 2019) (Figure 1C).

In addition to epithelial cells, Clusters 19, 21, and 27 were identified as fibroblasts with high expression of extracellular matrix (ECM) genes (COL3A1, COL1A1, POSTN, and DCN) (Figure 1C) (Han et al., 2020). Clusters 8 and 11 were annotated as smooth muscle cells (SMCs) with high expression of ACTC1, MYL4, and TTN (Han et al., 2020) (Figure 1C). Clusters 17, 18, and 22 were identified as proliferating cells with high expression of cell cycle-related genes, such as CENPF, TOP2A, and HIST2H2AC (Han et al., 2020) (Figure 1C). Cluster 26 cells were annotated as endothelial cells with high expression of VIM and PECAM1 (Han et al., 2020) (Figure 1C). Cluster 28 cells were identified as lymphocytes with high expression of ISG15, MX1, and DDX58 (Wani et al., 2019) (Figure 1C). Cluster 29 cells were identified as macrophages due to expression of C1QA, C1QB, and C1QC (Cochain et al., 2018) (Figure 1C), and Cluster 30 cells were identified as T cells expressing CD4 and CD69 (Schoenberger, 2012) (Figure 1C).

In summary, scRNA-seq successfully identified cell types of the rumen from the embryonic to functionally mature postnatal stages. This new comprehensive atlas of gene expression and marker genes in diverse cell clusters of the rumen provides a
Diverse cell-type subpopulations in the rumen

Given its importance in diet metabolism and microbial defense, rumen development and functional maturation play crucial roles in ruminant physiology (Baldwin & Connor, 2017). The rumen epithelium develops from two layers of cells at the embryonic state (i.e., stratum basale and stratum granulosum) to a four-layer structure after birth (i.e., stratum basale, stratum spinosum, stratum granulosum, and stratum corneum) (Garcia et al., 2012). Based on the identification of diverse epithelial cell types, we further investigated the genetic characteristics of the four strata of the rumen epithelium. Specific clusters were defined by cell-type abbreviation and characteristics of the four strata of the rumen epithelium.

Table 1  Summary of marker genes of each cell type identified in rumen

<table>
<thead>
<tr>
<th>Cell type</th>
<th>Subtype</th>
<th>Marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal cells</td>
<td>BAS_c0</td>
<td>KRT15, COL17A1, SYNE2</td>
</tr>
<tr>
<td>BAS_c6</td>
<td>IGFBP5, PDLIM4, PRD-SPRRII</td>
<td></td>
</tr>
<tr>
<td>BAS_c9</td>
<td>IGFBP6, IGFBP5, FABP5</td>
<td></td>
</tr>
<tr>
<td>Keratinocytes</td>
<td>JUN, FOS, ZFP36</td>
<td>–</td>
</tr>
<tr>
<td>DKs</td>
<td>FABP5, KRTDAP, SPINK5, CTA</td>
<td>DK_c2, DK_c4, DK_c10, DK_c12, DK_c23, DK_c24, DK_c25</td>
</tr>
<tr>
<td>DK_c2</td>
<td>FABP5, GSTA1, IGFBP2</td>
<td></td>
</tr>
<tr>
<td>DK_c4</td>
<td>CTA, IGFBP5, IGFBP6</td>
<td></td>
</tr>
<tr>
<td>DK_c10</td>
<td>RBP2, MGST1, TFF3</td>
<td></td>
</tr>
<tr>
<td>DK_c12</td>
<td>IGFBP2, IGFBP5, IDH1</td>
<td></td>
</tr>
<tr>
<td>DK_c23</td>
<td>MT-COX2, MT-CYT8, MT-ND4</td>
<td></td>
</tr>
<tr>
<td>DK_c24</td>
<td>P13, CSTA, LYPD1</td>
<td></td>
</tr>
<tr>
<td>DK_c25</td>
<td>MT-COX1, MT-ND4, MT-ATP6</td>
<td></td>
</tr>
<tr>
<td>TDKs</td>
<td>CNFN, KRT23, CAST</td>
<td>TDK_c14, TDK_c15, TDK_c16</td>
</tr>
<tr>
<td>TDK_c14</td>
<td>ANXA1, RPS2, RPL8</td>
<td></td>
</tr>
<tr>
<td>TDK_c15</td>
<td>AHNAK, DSP, JUP</td>
<td></td>
</tr>
<tr>
<td>TDK_c16</td>
<td>S100A9, CHMP2A, DSTN</td>
<td></td>
</tr>
<tr>
<td>SSCs</td>
<td>CA1, ACA2, BDH1</td>
<td>SSC_c1, SSC_c6, SSC_c13</td>
</tr>
<tr>
<td>SSC_c1</td>
<td>GSTA1, LYPD3, KRT17</td>
<td></td>
</tr>
<tr>
<td>SSC_c3</td>
<td>TST, CA1, YBX1</td>
<td></td>
</tr>
<tr>
<td>SSC_c5</td>
<td>DHRST7, CA12, KRT15</td>
<td></td>
</tr>
<tr>
<td>SSC_c6</td>
<td>MT-ND3, TFM1, IGFBP6</td>
<td></td>
</tr>
<tr>
<td>SSC_c13</td>
<td>PRD-SPRRII, S100A9, S100A12</td>
<td></td>
</tr>
<tr>
<td>Fibroblast cells</td>
<td>COL3A1, COL1A1, POSTN, DCN</td>
<td>Fibro_c19, Fibro_c21, Fibro_c27</td>
</tr>
<tr>
<td>Fibro_c19</td>
<td>MFAP5, DLK1, MGP</td>
<td></td>
</tr>
<tr>
<td>Fibro_c21</td>
<td>TXB3, NDF1, APOE</td>
<td></td>
</tr>
<tr>
<td>Fibro_c27</td>
<td>GPM6A, KRT8, LAMP5</td>
<td></td>
</tr>
<tr>
<td>Smooth muscle cells</td>
<td>ACTC1, MYL4, TTN</td>
<td>–</td>
</tr>
<tr>
<td>Endothelial cells</td>
<td>VIM, PECAM1, RAMP2</td>
<td>–</td>
</tr>
<tr>
<td>Proliferating cells</td>
<td>CENPF, TOP2A, HIST2H2AC</td>
<td>–</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>ISG15, MX1, DDX58</td>
<td>–</td>
</tr>
<tr>
<td>T cells</td>
<td>CD4, CD69, CD53</td>
<td>–</td>
</tr>
<tr>
<td>Macrophages</td>
<td>C1QA, C1QB, C1QC</td>
<td>–</td>
</tr>
</tbody>
</table>

*: Not available.
indicated that up-regulated DEGs in KERA_c20 cells were enriched in the tumor necrosis factor (TNF) signaling pathway (Supplementary Table S3), which is involved in cell survival, death, and differentiation (Webster & Vucic, 2020), thus suggesting that it may be crucial for keratinocyte differentiation. Comparison of DEGs (Supplementary Table S4) in the DK subpopulations showed that DK_c2 and DK_c10 cells highly expressed FABP5 and TFF3 (Figure 2B). FABP5 induces keratinocyte differentiation through regulation of fatty acid transport (Ogawa et al., 2011), while TFF3 is a modifying factor in pathways regulating cell proliferation and keratinocyte migration (Storesund et al., 2009). Therefore, these two cell types may be associated with keratinocyte proliferation and differentiation. The DK_c4 and DK_c24 cells highly expressed ANXA1 and CSTA (Figure 2B), which participate in the cornified envelope formation in keratinocytes (Ma & Ozers, 1996; Steven & Steinert, 1994), suggesting that these two subpopulations may be associated with keratinization. The three TDK subpopulations (TDK_c14, TDK_c15, and TDK_c16) showed different gene expression profiles (Figure 3A; Supplementary Table S5). TDK_c14 cells showed high expression of ribosomal protein genes. TDK_c15 cells showed high expression of AHNAK, DSP, and JUP, which are associated with desmoplasic plaques (Masunaga et al., 1995; Yang et al., 2006), indicating a contribution of TDK_c15 to cell adhesion and maintenance of ruminal epithelium stability. TDK_c16 cells showed high expression of innate immune-related genes (IRF7, ISG15, FADD, MAPK14, and MAPK13), suggesting that the stratum corneum participates in innate immunity in addition to physical defense.

The stratum spinosum is the main metabolic layer of the rumen and can be distinguished morphologically after birth (Baldwin & Connor, 2017). Comparison of the proportions of epithelial cells in different periods showed that five clusters (proliferating cell cluster PC_c17 and SSC clusters SSC_c1, SSC_c3, SSC_c5, and SSC_c13) were significantly more abundant at postnatal day 45 (Figure 1E). These five subpopulations showed high expression of key ketogenesis-related genes (ACAT1, SLC16A1, and BDH1) (Figure 3B; Supplementary Table S6). Ketogenesis is a hallmark of metabolic development of the rumen epithelium (Naeem et al., 2012). Therefore, we defined these five cell clusters as the
major cell types of the rumen epithelium stratum spinosum. Among the SSC subpopulations, SSC_c5 cells exhibited high expression of metabolic genes and marker genes of basal cells (KRT15 and IGFBP2) (Figure 2C), suggesting that they have a similar differentiation capacity to basal cells. SSC_c3 cells showed high expression of mitochondrial genes, suggesting a major role in energy metabolism. SSC_c1 and SSC_c13 cells showed high expression of SPINK5 and PRD-SPRRII, indicating that these two subpopulations exhibited the highest degree of keratinization. Interestingly, PC_c17 cells expressed more ketogenesis-related genes than other proliferating cells (PC_c18 and PC_c22), suggesting that PC_c17 may represent the proliferative state of SSCs (Figures 2C, 3B). Thus, we hypothesize that the SSC and PC_17 cell subpopulations may play key roles in the renewal and proliferation of the stratum spinosum, thereby maintaining the homeostasis of rumen metabolic function.

Fibroblasts, which are enriched in the rumen lamina propria, not only produce extracellular matrix (ECM) to provide structural support for the epithelium, but also participate in cell signaling, migration, proliferation, and differentiation (Kim et al., 2011). We identified three types of fibroblasts (Clusters 19, 21, and 27) in the rumen (Figure 2D). Both C19 and C27 cells showed high expression of COL14A1 (collagen-encoding gene) and MGP in comparison with C21 but could be distinguished by differential expression of MFAP5 and KRT8 (Figure 3C). C21 cells strongly expressed lipid transport-related genes APOE and APOD and collagen-encoding gene COL13A1 (Figure 3D). Comparison of the DEGs in the three fibroblast types indicated that C21 consisted of papillary fibroblasts, while C19 and C27 consisted of reticular fibroblasts. We further found that the papillary fibroblasts (C21) strongly expressed neuro-development (NDNF)-related genes, suggesting that papillary fibroblasts also participate in rumen innervation (Figure 3D).

In summary, combined with the characteristics of the rumen structure, we identified specific cell types associated with the morphological strata of the rumen and the heterogeneity of cells in these strata. Different cell subpopulations showed specific gene expression characteristics and different functions during rumen development.

Characterization of dynamic prenatal and postnatal rumen developmental patterns

Based on morphometric analysis (Figure 4A) and expression characteristics of samples obtained from seven periods of

![Figure 4](image-url)

**Figure 4** Characterization of rumen developmental patterns

A: Schematic of rumen development, as elucidated in this study, illustrating typical changes in morphology observed during rumen development. EL, epithelial layer; LP, lamina propria; ML, muscularis layer; SL serosa layer; RP, ruminal pillars. B: t-SNE plots of cell clusters, colored according to indicated periods of rumen development. C: Proportions of cells of indicated types at indicated sampling times. D: t-SNE plots of cell clusters at four developmental stages and four periods of middle-I stage.
rumen development (Figure 4B; Supplementary Table S7), we distinguished four major stages of development, i.e., early (0–30 days of gestation), middle-I (60, 90, 110, and 130 days of gestation), middle-II (newborn), and late (45 days old) (Figure 4D).

During the early stage of rumen development, the rumen becomes a small separate compartment of the primitive gastric tube. The ruminal wall consists of four layers: i.e., internal epithelial layer, middle pluripotential blastemic tissue layer, tunica muscularis (with an inner circular layer and outer longitudinal layer), and serosa (García et al., 2012; Jenkins & Tortora, 2016). Cell-type analysis showed that all cell types were present in the early stage, but SMCs were dominant (Figure 4C). We identified two SMC clusters (SMC_c8 and SMC_c11), which corresponded to the inner circular layer and outer longitudinal layer of the tunica muscularis (Martini, 2006). In addition, scRNA-seq confirmed the presence of immune cells (lymphocytes, macrophages, and T cells) in the early-stage rumen (Figure 4C), which has been controversial due to their small numbers and difficult detection by conventional methods.

In the middle-I stage of rumen development, the ruminal wall consisted of four layers: i.e., epithelium, lamina propria + submucosa, tunica muscularis, and serosa (Figure 4A). Morphological change include ruminal papillogenesis and maturation of the epithelial layer (Figure 4A). With the formation of the rumen epithelium, several epithelial cells (basal cells, keratinocytes, DKS, TDKS) became the dominant type in the rumen, and the proportions of all epithelial cell types increased significantly at 60 days of gestation (Figure 4C). We also observed significant changes in the proportions of various epithelial cell types with increasing gestation time (Figure 1E, 4C). The percentages of keratinocytes and TDKS decreased and increased with gestation time, respectively (Figure 4C), which has been controversial for a long time. We found that TDKS showed high expression of PI3 (Figure 5A), which is associated with elongation of the lateral cell membrane (Wang & Brieher, 2020), and a rapid increase in abundance in the early papillogenesis phase at prenatal day 90. We hypothesize that DK_c24 cells are a transitional state for keratinocytes that appear during early papillogenesis of the rumen, and P2 is the process of papillae keratinization.

To further elucidate the P2 keratinization process, we visualized dynamic gene expression patterns along the pseudo-time trajectory and classified them into four groups (Figure 5C; Supplementary Table S9). Group 1 contained 317 genes highly expressed in early keratinization, including SOX2, MEIS1, RIF1, APC, ID3, SMARCAD1, SMAD5, and FGFR2 (Figure 5E), which are associated with regulation of stem cell pluripotency (Kaneshia et al., 2021). These genes were enriched in the GO terms “nucleoplasm” and “DNA binding”, suggesting that cell differentiation is a major step in early keratinization (Supplementary Table S10). The main keratin-encoding gene in this group was KRT17 (Figure 5E). Group 2 included 364 genes highly expressed in the mid-stage of keratinization, including genes associated with mitochondrial formation, maintenance, and function. These genes were enriched in the “Ribosome” pathway, indicating high rates of protein synthesis (Supplementary Table S10). The main keratin-encoding gene expressed in this group was KRT15 (Figure 5E). Group 3 contained 306 genes highly expressed in the mid-stage of keratinization, including KRT4, ANXA7, VDAC1, and CNN3 (Figure 5E), which are associated with epithelial cell differentiation (Kaneshia et al., 2021). These genes were enriched in the GO terms “microtubule” and “focal adhesion” (Supplementary Table S10), suggesting
Figure 5 Molecular features of keratinization and papillogenesis during rumen development

A: UMAP plot of epithelial cells showing keratinization process at cellular level. B: Pseudo-time trajectory of P2 keratinization process and deconvolution of pseudo-time plot according to cell type. C: Heatmap showing four clusters of pseudo-time genes differentially expressed with P2 keratinization. D: Heatmap showing four clusters of pseudo-time genes differentially expressed with P1 keratinization. E: Expression levels of representative genes in cells ordered along trajectory (colored based on clusters to which they are assigned). F: Changes in expression patterns of TBX3 gene in papillary fibroblasts over time. Cells are color-coded with rumen samples. G: Schematic of papillary development. H: Violin plots showing relative expression of KRT36 in diverse epithelial cell subpopulations.
that cellular connections are enhanced during this stage of keratinization. Group 4 contained 1242 genes highly expressed in late keratinization and enriched in the KEGG terms “adhesion junction” and “bacterial invasion of epithelial cells” (Supplementary Table S10). Genes of the S100A family (S100A2, S100A4, S100A5, S100A9, S100A10, S100A12, and S100A13), which are associated with the immune system (Xia et al., 2018), were also highly expressed in the late stage (Figure 5E), indicating that the rumen epithelium develops defensive capacities in this stage.

The neatly arranged papillae are a striking morphological feature of the mature rumen, enlarging the surface area available for nutrient absorption and metabolism (Graham & Simmons, 2005). Anormogenesis of rumen papillae can trigger subacute ruminal acidosis and other nutritional metabolic diseases that can severely impair animal production and health (Underwood et al., 2015). Papillary fibroblasts, which are close to the epithelial layer and associated with papilla formation, showed higher TBX3 transcription factor expression than the reticular fibroblasts (Figure 5F). Previous studies have shown that abnormal expression of TBX3 in humans can cause papillary carcinoma of the skin, ulnar-mammary syndrome, and arrhythmia (Frank et al., 2012; Khan et al., 2020). High TBX3 expression can also promote expression of PI3 (Peres et al., 2015), which was strongly expressed in the papillogenesis-related DK_c24 cells. We speculate that the increased expression of TBX3 in the papillary fibroblasts may promote an increase in PI3 expression in the DK_c24 cells, leading to elongation of the lateral membrane of the epithelial cells and hence papillogenesis (Figure 5G). In addition, we found that the five cell clusters in the stratum spinosum highly expressed KRT36 (Figure 5H), a specific marker of filiform papillae (Brychtova et al., 2020). This suggests that stratum spinosum cells are in the papillae and corroborates the importance of rumen papillae as sites of energy metabolism.

Functional rumen epithelium formation after birth

With the introduction of a solid diet after birth, the proportion of SSCs (1, 3, 5, and 13) and PC_c17 cells increased and became dominant in the rumen epithelium and formed the stratum spinosum by postnatal day 45. We identified 407 genes highly expressed in SSCs. KEGG enrichment analysis of up-regulated DEGs revealed that 73 genes were enriched in metabolic pathways, including carbon metabolism, fatty acid metabolism, propanoate metabolism, and butanoate metabolism (Supplementary Tables S11, S12). These findings suggest that the major cellular components required for a functionally mature rumen epithelium are established before postnatal day 45.

As ruminants are exposed to a highly complex environment from birth, the rumen must not only develop physical defenses, but also an immune response. However, there were no significant differences in the expression levels of immune-related genes among the three types of immune cells in the rumen before and after birth (Figure 6A). Interestingly, we found that SSC_c13 was the most keratinized subpopulation of SSCs, and cells in this cluster showed high expression of immune-related genes (ISG15, IL36A, and S100A gene family) (Figure 6B). ISG15 plays a key role in the innate immune system, inducing natural killer cell proliferation (Perng & Lenschow, 2018). ISG15 also induces interferon (IFN) cytokines as an anti-mycobacterial response and is highly expressed in lymphocytes (Gao et al., 2020). Thus, we hypothesize that SSC_c13 cells induce lymphocyte proliferation to generate an immune response. These findings suggest that maturation of the postnatal rumen stratum spinosum is not only important for ruminant energy

Figure 6 Molecular features of various cell types in rumen of lambs after birth

A: Violin plots showing expression of marker genes of three immune cell types at different developmental stages. B: Relative expression levels of immune-related genes in SSCs.
metabolism, but also for immune capacity.

CONCLUSIONS
As an important evolutionary innovation and unique organ, the rumen has played an important role in the adaptation of ruminants to their complex ecological environments. This study provides a detailed account of gene expression during prenatal to postnatal rumen development based on scRNA-seq analysis, including detailed characterization of the cell types involved, as well as identification of additional marker genes for specific cell types. We revealed the dynamic changes and high heterogeneity of the cellular and molecular profiles during four development stages of the rumen, providing clues for maternal regulation of fetal rumen development. However, this study is limited by the small sample size and number of captured cells.

The rumen has evolved a series of distinct structural hallmarks, such as neatly arranged papillae and keratinized stratified squamous epithelium, as well as innovative functional characteristics, such as VFA metabolism and immunology. Our identification of diverse basal cells, keratinocytes, DKs, TDKs, and SSCs provides important insight into rumen structure and function and a basis for isolating different cells in the rumen. The epithelial cells exhibit two processes during papillary keratinization accompanied by dynamic changes in identified gene expression. In addition, the cellular and gene expression profiles strongly suggested that the rumen epithelium is functionally mature by postnatal day 45, and the enriched stratum spinosum cells play crucial roles in VFA energy metabolism and immune capacity. Importantly, we identified the growth factor TBox3 as a potential marker gene for papillary growth by interacting with P13, thus highlighting the need to validate the role of TBox3 in papillogenesis in vivo and in vitro.

Our findings provide cellular-level insights into the molecular characteristics of morphogenesis, notably keratinization and papilla formation, and functional establishment (metabolism and immunity) of the rumen. This study will contribute to our understanding of rumen development and the interactions between dietary metabolism and the rumen.

DATA AVAILABILITY
The single-cell RNA-sequencing data were deposited in the Genome Sequence Archive database (http://gsa.big.ac.cn/) under accession No. CRA007511, Science Data Bank (doi:10.577 60/sciedb.j00139.00007), and NCBI under BioProjectID PRJNA857336.

SUPPLEMENTARY DATA
Supplementary data to this article can be found online.

COMPETING INTERESTS
The authors declare that they have no competing interests.

AUTHORS’ CONTRIBUTIONS
Q.Q., J.H.L., Z.P.L., and S.Y.M. designed the project and research; D.M.S., J.H.L., W.Y.Z., Y.Y.Y., and J.H. collected samples and constructed sequencing libraries; Y.Y., D.M.S., and T.Q. conducted single-cell data analysis; Q.Q., J.H.L., Z.P.L., and Y.Y. wrote the manuscript. Q.Q., J.H.L., Z.P.L., R.H., and Y.Y. amended the manuscript. All authors read and approved the final version of the manuscript.

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