Molecule-based osteoarthritis diagnosis comes of age

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Background

Osteoarthritis (OA) is a common disabling condition worldwide, representing a substantial and growing health burden with extensive socio-economic costs (1). Patients with OA typically manifest several symptoms that impair their quality of life, such as pain, stiffness, and dysfunction. Even though OA has been recognized as early as the late eighteenth century (2), the current diagnosis of it is largely dependent on clinical information; including symptoms, signs, and images. Such diagnostic criteria are not able to predict high risk OA individuals and/or provide evidence for early diagnosis of the disease. In addition, despite having emerging pharmaceutical therapies in recent decades (3), only a fraction of potentially disease-modifying OA drugs have been applied in clinical practice. Whereas, a portion of these drugs show ambiguous outcomes, and the use of them in clinical guidelines are usually in disagreement (4,5). This could be due to a mismatch between the molecular mechanisms by which the drug works and the clinical manifestations at the time of the decision to use the drug, since the commonly recognized clinical features cannot elucidate the pathological changes of OA (6). Thus, the lack of pathophysiology-based OA diagnosis impedes the development of targeted therapeutics.

OA is a molecular disorder

OA is a disease of the whole joint involving the structural changes in the articular cartilage, synovium, subchondral bone, ligaments, and periaricular muscles (1). These structural changes are the cumulative results of changes at the molecular level, as the definition of OA proposed by the Osteoarthritis Research Society International: “The disease manifests first as a molecular derangement (abnormal joint tissue metabolism) followed by anatomic, and/or physiologic derangements (characterized by cartilage degradation, bone remodeling, osteophyte formation, joint inflammation and loss of normal joint function), that can culminate in illness” (7). For instance, under pathological conditions, the depletion of cartilage composition, mainly proteoglycan and collagen, leads to a loss of cartilage integrity and a change in the properties of cartilage material (8). These changes increase the susceptibility to cartilage rupture and eventually result in cartilage degeneration, which can be seen on magnetic resonance imaging (MRI) as thinning of the cartilage or on X-ray as narrowing of the joint space (9) during later stages in OA. Therefore, detection of early altered molecules seems to provide opportunity for a more sensitive and accurate pathological diagnosis of OA.

Transcriptome atlas-based OA diagnosis

In this context, research in the field of molecular diagnosis of OA has been very active in recent years. Since the pattern of gene expression reflects cell responses to pathological condition, transcriptome data has been used in the diagnosis and classification of OA in several studies. For example, Soul et al. have investigated the pattern of gene expression in non-OA and OA cartilage by RNA-sequencing (RNA-seq) (10). To avoid the gene expression alterations that occur in damaged cartilage, they analyzed the intact cartilage in OA group. They showed 2,692 differentially
expressed genes between non-OA and OA cartilage, and surprisingly, they found a large increase in the expression of matrix protein genes in the OA group. Further unsupervised clustering analysis stratified OA into two subgroups: Group A showed increased expression of cartilage components like collagen type II, V, IX and XI and less expression of collagen type I; in contrast, Group B showed reduced expression of chondrogenic genes and enhanced expression of osteogenic genes. Similarly, Coutinho de Almeida et al. analyzed the whole-transcriptome profiling of OA cartilage and also identified two subgroups (11). Upon integrating radiographic OA data, they found that one subgroup was likely to be characterized by lower osteophyte scores and higher joint space narrowing (JSN) scores. These results clearly show that OA is a highly heterogeneous disease and raises the concept for a more precise molecular diagnosis.

Another example of an unexpected significant discovery was recently published in Bone Research. Different from previous studies that only focused on cartilage, Yuan et al. constructed an extensive transcriptome atlas of OA cartilage, subchondral bone and synovium, which were mainly affected tissues during OA pathophysiology (12). Through applying the unsupervised clustering analysis, they classified OA into four subtypes: cluster 1 (C1) subtype with glycosaminoglycan metabolic disorder, C2 subtype with collagen metabolic disorder, C3 subtype with activated sensory neuron, and C4 subtype with inflammation. They further linked the clinical symptoms of different OA subtypes with molecular functions by the ligand-receptor crosstalk analysis of cartilage, subchondral bone and synovium. For example, they found more tissue crosstalk in C4 subtype than others, especially ossification-enriched subchondral bone-cartilage crosstalk and osteoblast differentiation-enriched subchondral bone-subchondral bone crosstalk, suggesting subchondral bone overgrowth and narrowed joint space may be observed in C4. Further, upon clinical data analysis for each OA subtype, they found that C4 patients had a higher JSN score, demonstrating JSN might be the trait of C4 OA patients. This study is surely of great importance as it links the pathogenesis of OA patients with their clinical manifestations. Their findings provide a new approach for the diagnosis of OA, and the transcriptome atlas may allow for precise diagnosis and targeted therapeutics of OA in the future.

In recent years, RNA-seq (10-12) and single-cell RNA-seq (13,14) have been extensively studied in order to reveal OA subtypes and pathophysiology at molecular and cellular levels. However, it should be critically noted that the transcriptome-based diagnosis has some limitations. First, although transcriptional networks play a fundamental role in governing cell function and fate, they do not entirely determine cellular identity due to ubiquitous post-transcriptional regulation, translational regulation, and degradation mechanisms (15). Confirmation of the link among transcriptome atlas, proteome atlas, and clinical information for OA patients is still pending. Second, the aforementioned transcriptome analysis relies heavily on the technically and invasively acquired joint tissue specimens. It is difficult to generalize in clinical practice, not available for early OA diagnosis and cannot dynamically reflect the pathological changes of the disease, especially for post-treatment evaluation.

Body fluid: readily accessible molecular pool for OA diagnosis

One of the most important contributors of the OA diagnostic algorithm shifts to a molecule-based assessment is the research on potential biomarkers. According to the World Health Organization definition, a biomarker is “any substance, structure, or process that can be measured in the body or its products and influence or predict the incidence of outcome or disease” (16).

Numerous biomarkers that reflect the pathophysiology of OA can be found in body fluids, including synovial fluid (SF), blood, and urine. SF represents the gold standard fluid for the identification of biomarkers in OA because of its intimate relationship with various joint tissues. Changes in tissue environment will directly affect the SF composition. There is extensive evidence that the level of cartilage oligomeric matrix protein (COMP) in SF is significantly increased after partaking in a marathon (17), suggesting that COMP in SF may be a sensitive indicator of cartilage wear. Blood is easily withdrawn and mediates many of the immunologic pathways. The most widely studied OA biomarkers through blood test are inflammatory cytokines like interleukin-1β, tumor necrosis factor-α, and C-reactive protein, which are highly associated with symptomatic and radiographic progression of OA (18,19). Their levels may provide the possibility to dynamically monitor the severity and progression of OA. In addition, urine is easily accessible with large volumes and can be acquired with noninvasive techniques. Some metabolites are detectable in urine, like C-telopeptide fragments of Collagen II (CTX-II) and C-telopeptide of Collagen I, which are two recognized indicators of collagen degradation and are associated with
OA progression (20,21). Of note, the urinary levels of CTX-II are responsive to chondroprotective glucosamine (22). In this context, the evaluation of urinary CTX-II levels may allow for the stratification of OA patients who may benefit most from chondroprotective therapeutics and help to monitor the treatment efficacy dynamically.

Taken together, the molecules in body fluids could provide us with lots of information about OA pathogenesis, which will greatly promote the development of molecular diagnosis of OA. By associating molecular profiles in body fluids with clinical information, OA stages and the efficacy of some therapeutics, a more comprehensive and scientific picture should be created to test the clinical application value of molecular diagnosis of OA.

**Future perspectives**

In conclusion, the excellent researches mentioned above bring to the light an important issue concerning molecular diagnosis of OA, providing a novel approach for the definition of disease subtypes. Since OA is a highly heterogeneous disease, molecule-based diagnosis raises the possibility that different subgroups may be adapted to different modes of intervention. Therefore, molecular characteristics should be considered in the development of inclusion and exclusion criteria in future clinical trials. Stratifying homogenous patients at the molecular level and selecting therapies targeting their pathogenesis may potentiate the efficacy of some therapeutics. We hope the following studies will focus on deepening the knowledge of molecular profile during OA initiation and progression, and validating their relevance to clinical practice, which will contribute to further improvements in therapeutic options for patients with OA. In the future, biomarkers in body fluids together with clinical information may become the first screening method for OA diagnosis and classification, and transcriptomic and/or proteomic analysis of joint tissues may be effective tools for validating OA subtypes.

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