Angiotensin I–converting enzyme (ACE, or DCP1) is a zinc metallopeptidase that converts angiotensin I into the vasoactive and aldosterone-stimulating peptide angiotensin II and cleaves bradykinin into inactive peptides. Plasma ACE measurement is widely used for the diagnosis of sarcoidosis. While enzyme concentrations are highly stable in an individual, there is a high level of interindividual variability. In 1990, we identified an insertion/deletion polymorphism in ACE that functions as a quantitative trait locus (QTL), accounting for half of the interindividual variability. Since then, technological advances have allowed for the elucidation of expression QTLs (eQTL). Such studies are allowing researchers to determine how underlying genetic predisposition contributes to human disease.

After we determined the amino acid structure of ACE by cloning its cDNA, we used the ACE cDNA probe as a genetic tool for exploring both the ACE plasma level polymorphism and the potential role of the gene in the predisposition to cardiovascular and renal diseases. The JCI article by Rigat et al. that we comment on here showed a clear relationship between an ACE marker genotype consisting of two different alleles, the insertion (I) and the deletion (D) allele within intron 16 of the gene, and the serum level of the enzyme, which is known to be stable within a given individual, but to differ among individuals (6). A previous segregation study in healthy families by Cambien et al. showed that ACE serum levels were under the influence of a “major gene,” also called a quantitative trait locus (QTL) (7). Using a DNA probe for the ACE gene (8), we identified the genetic locus underpinning the observed QTL, demonstrating that an ACE gene polymorphism was acting in cis on the serum ACE level, as well as on the ACE mRNA level (9). This strong relationship has been replicated several times since.

Since the polymorphic insertion is a repetitive element within an intron of ACE, we were keen to determine whether this I/D polymorphism was itself responsible for the expression variation in ACE or whether it was only a marker in linkage disequilibrium with functional variants. Using additional ACE markers in 96 families, we showed that there were indeed two QTL within the ACE gene, one located in the S′ region, and another one in the central part of the gene (10). These results were confirmed in a later study, which included an extensive search for ACE gene variants, with subsequent genotyping of these variants in a larger series of 250 healthy families of mixed European descent. Clade analysis of the haplotypes was then performed in order to search for associations with ACE serum levels (11). Two recombination break points were detected in the major haplotypes, one upstream of the transcription initiation site and the other in the 3′ region flanking the ACE gene. The major effect was due to variants located between these two recombination points, and a weaker effect was linked to the upstream region. Indeed, the high-linkage disequilibrium observed between the markers in European subjects prevented further identification of the functional variant. Greater haplotype diversity was observed in West African families, resulting in the exclusion of the I/D polymorphism as the functional variant. These results indicated that multiple sites influence circulating ACE levels, among which four potential sites were candidates for a direct effect (12). Further in vitro studies with cloned gene fragments could not confirm the functional role of these variants (our unpublished results).

Thus, 20 years after its initial publication, the molecular basis underlying the expression QTL (eQTL) of the ACE gene has not been elucidated at the DNA level.

Such a relationship involving a major RAS gene spurred numerous association studies focused on discovering the links between the ACE eQTL and a genetic risk for cardiovascular or renal diseases. Results of case-control studies are often conflicting and prone to errors due to recruitment bias, population stratification, and multiple comparisons, requiring validation in independent groups of subjects. The hundreds of studies that have taken place since the identification of the ACE I/D polymorphism have all aimed to discover associations between the ACE I/D marker and various diseases involving the renin angiotensin or kal-
likrein kinin systems, such as myocardial infarction or coronary heart disease, stroke, renal diseases, or complications of diabetes (13–17). Study results were generally conflicting, but a recent meta-analysis of 50 case-control studies on the ACE I/D polymorphism and stroke concluded that carrying the ACE D allele constituted a low penetrance risk factor (18). Indeed, meta-analyses conducted with rigorous statistical methodology will help to clarify any associated risk with the ACE gene polymorphism, although a bias toward the publication of positive results is possible.

Genetically engineered mice carrying either a single copy, or two or three active copies of the ACE gene were generated in the Smithies laboratory, allowing the effects of increased ACE expression to be tested in different pathological models (19). Indeed, three-copy mice presented with increased proteinuria after 12 weeks of diabetes, as compared with mice carrying only one or two copies (20). ACE mRNA was increased in the kidney of three-copy mice, and decreased bradykinin levels were proposed to account for the marked effects on glomerular hemodynamics during diabetes in these mice. This experimental model combining genetically increased ACE and diabetes represents a valuable support for the effect of the ACE I/D polymorphism on diabetes complication in humans (21).

The ACE gene represents a paradigm of eQTL, and within 20 years, a complete revolution has occurred. eQTL searches have benefited from the “omics” revolution, combining genome-wide SNP genotyping, transcript expression analysis via microarray or RNAseq, and determination of thousands of protein levels in biological fluids by proteomic analysis (22). Combining these measurements allows a genome-wide search for eQTL that are relevant to disease susceptibility, such as the study performed by Chambers et al. for the plasma concentration of liver enzymes (23). These eQTL are essential for understanding the associations between SNPs and complex diseases by genome-wide association studies. By identifying a growing number of these relationships, we will be able to put together, like pieces of a puzzle, the elementary traits underlying genetic predisposition to cardiovascular diseases.

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