REVIEW ARTICLE

Molecular methods for assessing insect parasitism

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Abstract

Determining insect parasitism rates is problematic due to the small size and lack of useful distinguishing morphological characters of many parasitoid taxa. To solve this problem, entomologists have employed one of four general methods to detect parasitoid protein or nucleic acid markers: serological assay; random amplified polymorphic DNA-polymerase chain reaction (RAPD-PCR); allozyme electrophoresis; or specific PCR. Serological methods, especially with monoclonal antibodies, are unrivalled for specificity, enabling discrimination at the stage as well as species level. However, they have not found favour with many workers, possibly due to complexity and expense. RAPD-PCR has been widely used, but can only be recommended for restricted applications because of its poor reproducibility. Allozyme electrophoresis provides reproducible detection and discrimination of closely related species. Specific-PCR is highly specific and reproducible, and also has the shortest latency for detection, usually 24 h or less after parasitization. The substantial existing literature on allozyme electrophoresis and specific PCR is used to support recommendations on what are apt to be fruitful enzyme systems or genomic regions for detecting and discriminating parasitoids in untried parasitoid-host assemblages.

Keywords: allozyme electrophoresis, isozyme, specific PCR, parasitoids, RAPD-PCR

Introduction

Parasitioids are parasites that kill their hosts as a normal part of their development (Lafferty & Kuris, 2002). This functional group comprises a broad taxonomic array of both parasitoid and host organisms, e.g. flagellates attacking diatoms (Kühn & Hoffmann, 1999) and flatworms attacking decapod crustaceans (Kuris *et al.*, 2002). To practising entomologists, however, parasitoids are insects, primarily in the orders Hymenoptera and Diptera, whose larvae develop in or upon the bodies of other insects. They destroy billions of pest insects annually in crops worldwide and are key players in biological control. While most parasitism is

*Fax: 301 504 5104 E-mail: greenstm@ba.ars.usda.gov ambient, parasitoids are also produced in vast quantities for intentional inundative release to control pest infestations (Pinto *et al.*, 1993; Ram *et al.*, 1995; Burks & Pinto, 2002; Borghuis *et al.*, 2004). Numerous parasitoid species have also been released and established on new continents to rejoin their hosts in what had been 'enemy-free space' (Jeffries & Lawton, 1984; Loxdale & Lushai, 1999), in classical biological control programmes to suppress exotic invasive pest species (Takada, 1998; Kimani-Njogu *et al.*, 1998; Hufbauer *et al.*, 2001; Prinsloo *et al.*, 2002; Iline & Phillips, 2004; Persad *et al.*, 2004).

Understanding and managing parasitoid-host interactions pose daunting technical challenges. Most parasitoids are small, even minute, and many groups are speciose and evolving rapidly. Related species often have few or no known sufficiently invariant distinguishing morphological characters for reliable discrimination (Pungerl, 1986; Landry *et al.*, 1993; Pinto *et al.*, 1993; Demichelis & Manino, 1998; Kimani-Njogu et al., 1998; Stouthamer et al., 1999; Barnay et al., 2001; Chang et al., 2001). Detecting and differentiating endoparasitic stages within their hosts is particularly difficult. Parasitoids have been visualized within hosts by conventional X-ray (Biever & Boldt, 1970) and magnetic resonance imaging (MRI; Geogehegan et al., 2000). While these techniques might enable detection of parasitism per se, current resolution would be insufficient to discriminate among closely related parasitoid species (Hart et al., 2003), if indeed morphological differences could be found among the immature stages of related species (Persad et al., 2004). Another visualization technique, near-infrared spectroscopy, has been used to separate adults, but not immatures, of sibling species (Cole et al., 2003). Hagler & Jackson (1998) used exogenous molecules, vertebrate immunoglobulins, to mark adult parasitoids for later identification.

Most biological control workers employ one of two methods for detection and discrimination of parasitoids: holding field-collected hosts in the laboratory for parasitoid emergence, or dissecting hosts and examining endoparasitic stages microscopically. Holding for emergence requires provisioning of artificial diet or living plant material to sustain the hosts until parasitoid emergence, usually under temperature-controlled conditions. Because of the time lag between host collection and parasitoid emergence, the results become available too late to be of use for management decisions. In the worst case, some species may have a prolonged post-emergence diapause (Keen et al., 2001), sometimes approaching a full year (Tilmon et al., 2000). Even when this is not a problem, there is almost always a failure of some hosts to produce parasitoids, sometimes due to patent microbial infections (Greenstone & Edwards, 1998; Ratcliffe et al., 2002), but often due to unknown causes, and for the latter there is no way to determine the status of these so-called 'duds' (Wool et al., 1978; Stuart & Greenstone, 1996; Ratcliffe et al., 2002). Whatever the cause, this leads to unknown biases in percentage parasitism estimates. Dissection on the other hand does provide unequivocal and timely estimates of parasitization rate, but it is tedious and requires special training, and species-specific identification can be challenging where members of the same genus or even family are present (e.g. Persad et al., 2004).

This has led to a search for stable, unequivocal, speciesspecific characters to support evolutionary, ecological, and applied research. Such characters are generally drawn from two classes of organic molecules: proteins and nucleic acids. These have the virtue of abundance in tissues, ubiquity within animals in general, and, in many cases, assignability to loci inherited in simple Mendelian fashion. Furthermore, formats for sensitive and reproducible assays of these molecules are available. This has made it possible to use them for studies of phylogeny (Campbell et al., 1993; Kimani-Njogu et al., 1998), genetic variability (Landry et al., 1993; Baldanza et al., 2001; Baker et al., 2003; Chaterjee et al., 2003; Baer et al., 2004; Hufbauer et al., 2004), hybridization (Atanassova et al., 1998), population dynamics (Vaughn & Antolin, 1998; Schneider et al., 2003; Macdonald & Loxdale, 2004), effectiveness of release and establishment (Edwards & Hoy, 1995; Prinsloo et al., 2002; Ratcliffe et al., 2002) and quality control (Yazlovetskii et al., 1981; Unruh et al., 1983; Morgan et al., 1988; Stuart & Burkholder, 1991; Landry et al., 1993; Roehrdanz et al., 1993), as well as for our present purposes, differentiation and detection within the host.

With the large literature presently available, it is possible to gain an impression of which proteins and which DNA regions are potentially most useful for detecting and discriminating parasitoids. Here I summarize this literature to determine which molecules represent sufficiently variable and detectable characters for parasitoid detection and discrimination; I then provide some guidelines on which specific characters appear most useful for these tasks.

Proteins

Animal tissues contain an abundance and diversity of enzymes and structural proteins. Techniques for detecting and characterizing both protein classes are well established and both have been used to discriminate and detect parasitoids.

Enzymes

Enzymes may be found as multiple forms, produced by different loci, referred to as isozymes; when there is more than one allele at an isozyme locus, each is referred to as an allozyme (Richardson *et al.*, 1986). These different enzyme forms usually migrate to different positions on electrophoresis gels and are visualized as bands that become stained when exposed to appropriate substrates and linking systems (enzymes, coenzymes); if they do not differ in mobility, they can often be differentiated by means of specific stains (e.g. Wynne *et al.*, 1992). The genetics, chemistry, and interpretation of allozyme electrophoresis are thoroughly explained by Richardson *et al.* (1986). The mobility and number of bands may be influenced by a number of factors, including pH, viscosity of the matrix, and ionic strength of the electrophoresis buffer.

Allozymes by definition reveal simple Mendelian traits (Kimani-Njogu et al., 1998), permitting genetic analysis (Atanassova et al., 1998), and most practitioners assume that electrophoretic banding patterns reflect such inheritance. In fact this is not always the case, because the number of alleles may not be revealed due to non-detectable amino acid substitutions (Richardson et al., 1986). Bands may also differ in staining intensity due to differences in allele number (homozygosity versus heterozygosity), subunit structure of enzymes affecting expression of bands in homozygous and heterozygous condition (Richardson et al., 1986), and gene duplication. Post-transcriptional modifications of the polypeptide chains can also complicate the interpretation of banding patterns (e.g. Cameron et al., 1984). If the assumption of Mendelian inheritance is critical to the analysis, formal genetic crosses should be made to confirm it (for examples, see Hung & Huo, 1985, and Pinto et al., 1992).

Because Mendelian inheritance cannot be assumed, the most conservative authors refer to electrophoretic variants as electromorphs rather than alleles. For our purposes, these bands will be useful as diagnostic markers provided they confer the specificity and sensitivity required for detection and discrimination, regardless of the mode of inheritance. Whether particular electromorphs are reliable species identifiers, i.e. invariant within species but differing between species, must be determined empirically. This can produce some surprises. For example, host switching may lead to loss of esterase bands (Cameron *et al.*, 1984), indicating that allozyme expression is subject to rapid evolution.

Scores of enzymes have been used in animal isozyme studies (Richardson *et al.*, 1986). Numerous authors have assayed a dozen or more enzyme systems in their search for sufficiently polymorphic bands/loci to enable discrimination of parasitoid species within particular parasitoid–host systems (Castrovillo & Stock, 1981; Castañera *et al.*, 1983; Pinto *et al.*, 1992; Atanassova *et al.*, 1998; Kimani-Njogu *et al.*, 1998; Burks & Pinto, 2002). Wynne *et al.* (1992) provide a list of eight enzymes they have found to be generally polymorphic in insects.

Table 1 provides an overview of enzyme systems that have been used as markers in all manner of parasitoid research. In each case, the table identifies those that the authors found most useful for differentiating parasitoids in the particular host–parasitoid systems under study. Not surprisingly, allozymes of hyperparasites as well as primary parastoids are detectable and may be species-specific (Cameron *et al.*, 1984).

There have been relatively few assessments of the temporal threshold for detection of allozymes of parasitoids within their hosts. Castañera *et al.* (1983) were able to detect 'nearly full grown' larvae of *Aphidius uzbekistanus* (Luzhetski) (Hymenoptera: Aphidiidae) in *Sitobion avenae* (Fabricius) (Hemiptera: Aphididae). Walton *et al.* (1990a) detected parasitism of *S. avenae* after 84 h by *Praon volucre* (Haliday) (Hymenoptera: Braconidae) (24 h later than by dissection); after 156 h by *Ephedrus plagiator* Nees (Hymenoptera: Aphidiidae) (72 h later than by dissection); and after 132 h in *Aphidius rhopalosiphi* De Stefani Perez (Hymenoptera: Aphidiidae) (24 h earlier than by dissection).

Structural proteins

Proteins may be detected and characterized by a number of standard biochemical techniques, e.g. native and denaturing polyacrylamide gel electrophoresis (PAGE and SDS-PAGE, respectively), and isoelectric focusing (IEF). SDS-PAGE has been used to differentiate closely related parasitoid species (Baldanza *et al.*, 2001). However, the most useful approach for parasitoid differentiation and detection has been immunoassay.

The vertebrate immune system, which is capable of recognizing proteins and protein substructures with exquisite specificity (Greenstone, 1996), can provide antibodies as reagents for use in standardized assays. Recognition of the parasitoid antigen protein by the antibody is made visible by an enzyme-substrate reporter system, in an immunodot assay or enzyme-linked immunosorbent assay (ELISA).

Within an immunized vertebrate, antibodies are produced by clones of lymphocytes, each clone secreting into the blood stream a population of identical antibody molecules recognizing a single region of an antigenic molecule, the so-called antigenic determinant (or epitope). Since proteins typically contain more than one determinant, more than one lymphocyte clone will be activated and the resulting polyclonal antiserum will recognize numerous determinants. Therefore when antiserum from an immunized animal is used as the reagent in a parasitoid assay, the assay is apt to cross react (produce false positives) with related species due to the presence of shared antigenic determinants, a problem that may not be solved by absorption with specific antigen (e.g. Keen et al., 2001). The surest route to avoiding cross-reactivity is the use of monoclonal antibodies, which are produced by lymphocyte

clones removed from the immunized vertebrate host, immortalized by fusion with a myeloma cell, isolated in tissue culture, and selected to secrete antibodies that recognize only unique determinants (Greenstone, 1996). Since determinants vary in their distribution across taxa, monoclonals may be specific at almost any taxonomic level, and even different developmental stages of a single species. For example, Stuart & Greenstone (1996) developed a monoclonal that distinguished first from second instars of just one braconid species, and another (Stuart & Greenstone, 1997) that recognizes all hymenopterous parasitoids tested, but not their hosts. This was due to the determinant being a hymenopteran-specific variant of elongation factor-1 α (Stuart, 1998), a protein common to all animals.

In addition to their own structural proteins, parasitoids may also secrete specific proteins into their hosts (e.g. Soldevila & Jones, 1991; Hochuli & Lanzrein, 2001). Although these do not appear to have been used as markers of parasitization, they should in principle work as well as parasitoid structural proteins.

Immunoassays are widely used for diagnosis but have not been much used to document parasitoid-host interactions. Consequently there are few data on the time course of detection of parasitoid antigens within hosts. A polyclonal ELISA for a pteromalid parasitoid (Hymenoptera: Pteromalidae) of house fly pupae (*Musca domestica* Linnaeus (Diptera: Muscidae)) could not detect larvae until 7 days postparasitization (Keen *et al.*, 2001), and another for a braconid wasp could not detect 'early stage' larvae (Allen *et al.*, 1992) in its lepidopteran leaf roller host. A monoclonalbased immunodot assay of Stuart & Greenstone (1996) was able to detect first instars of a braconid in its caterpillar host.

Nucleic acids

An advantage of nucleic acids is the large number of loci revealed, far in excess of those available for protein analysis (Edwards & Hoy, 1993; Landry *et al.*, 1993). Although ribonucleic acid (RNA) could in principle be used and would enable stage- as well as species-specificity, only deoxyribonucleic acid (DNA) has been employed to date. Thanks to the ability of the polymerase chain reaction (PCR; Ehrlich, 1989) to amplify minute quantities of DNA, scores of reactions can be performed on a single individual. Protein detection generally requires a larger quantity of target, so that fewer reactions can be run on a single animal (Edwards & Hoy, 1993; Vanlerberghe & Chavigny, 1997).

The sensitivity of a molecular assay depends upon the number of copies of the target molecule in the sample. DNA targets may be either non-coding sequences or specific genes. Multiple copies of a given sequence are found throughout the genome, variously arrayed among chromosomes in the nucleus or within cellular organelles. Both within and among individual animals and populations, these sequences may be identical or variable to a greater or lesser degree; they also display varying rates of evolution (Hillis & Dixon, 1991; Elder & Turner, 1995; Hoy, 2003), suggesting that some are better suited than others for species-level discrimination. Nevertheless, all of these classes of sequences have been successfully used to discriminate and detect closely related parasitoid species.

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Enzyme	Application	References
Acid phosphatase	Discrimination	Burks & Pinto, 2002 Pinto <i>et al.</i> , 1993
Aconitase	Discrimination	Pinto <i>et al.</i> , 1993 Unruh <i>et al.</i> , 1983
Esterases	Discrimination and detection Discrimination	Castrovillo & Stock, 1981 Burks & Pinto, 2002 Castañera <i>et al.</i> , 1983 Dawah, 1989 Dawah <i>et al.</i> , 2002 Cameron <i>et al.</i> , 1094 Kimani-Njogu <i>et al.</i> , 1998 Silva <i>et al.</i> , 1999 Pintureau, 1993 Ram <i>et al.</i> , 1995 Pungerl, 1986 Ram <i>et al.</i> , 1995 Silva <i>et al.</i> , 1999 Takada, 1998 Unruh <i>et al.</i> , 1983 Yazlovetskii <i>et al.</i> , 1981
	Discrimination and detection	Castrovillo & Stock, 1981 Walton <i>et al.</i> , 1990a,b Yazlovetskii <i>et al.</i> , 1981
Calcium binding protein	Discrimination	Iline & Phillips, 2004
Creatine kinase	Discrimination	Morgan et al., 1988
Fumarase	Discrimination	Pinto <i>et al.,</i> 1993 Unruh <i>et al.,</i> 1983
Glucose-phosphate isomerase	Discrimination	Burks & Pinto, 2002 Kimani-Njogu <i>et al.</i> , 1998 Kitthawee <i>et al.</i> , 1999 Pinto <i>et al.</i> , 1993
Glucose-6-phosphate dehydrogenase	Differentiation	Burks & Pinto, 2002 Pinto <i>et al.</i> , 1993
Glutamate-oxaloacetate transaminase	Discrimination and detection	Walton <i>et al.,</i> 1990a Dawah <i>et al.,</i> 2002
Glyceraldehyde-3-phosphate dehydrogenase α -glycerol-phosphate dehydrogenase	Discrimination Discrimination	Morgan <i>et al.,</i> 1988 Burks & Pinto, 2002 Morgan <i>et al.,</i> 1988
Glycerol-3-phosphate dehydrogenase Hexokinase	Discrimination and detection Discrimination Discrimination	Demichelis & Manino, 19 Kitthawee <i>et al.</i> , 1999 Dawah, 1989 Kimani Nigay <i>et al.</i> , 1908
Sorbital dehydrogenase	Discrimination	Kimani-Njogu et al., 1998 Kimani-Njogu et al., 1998
Isocitrate dehydrogenase	Discrimination	Atanassova <i>et al.</i> , 1996 Atanassova <i>et al.</i> , 1998 Burks & Pinto, 2002 Kitthawee <i>et al.</i> , 1999 Morgan <i>et al.</i> , 1988 Pinto <i>et al.</i> , 1993 Unruh <i>et al.</i> , 1983
Lactate dehydrogenase Leucine aminopeptidase Dihydrolipoamide dehydrogenase Malate dehydrogenase	Discrimination and detection Discrimination and detection Discrimination Discrimination	May et al., 1977 Castrovillo & Stock, 1981 Iline & Phillips, 2004 Iline & Phillips, 2004 Kitthawee et al., 1999 Morgan et al., 1988 Pinto et al., 1993 Tomiuk et al., 1979 Wool et al., 1978
	Detection	Castañera et al., 1983
Malic enzyme	Discrimination and detection Discrimination	Walton <i>et al.</i> , 1990a Castañera <i>et al.</i> , 1983 Unruh <i>et al.</i> , 1983 Walton <i>et al.</i> , 1990a Burks & Pinto, 2002 Pinto <i>et al.</i> , 1993 Pungerl, 1986 Kitthawee <i>et al.</i> , 1999

Table 1. Most useful enzyme systems used in parasitoid allozyme electrophoresis.

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Enzyme	Application	References
Mannose phosphate isomerase	Discrimination	Kitthawee et al., 1999
Peptidase	Discrimination	Atanassova et al., 1998
Phosphoglucomutase	Discrimination	Atanassova et al., 1998
1 0		Morgan et al., 1988
		Burks & Pinto, 2002
		Dawah, 1989
		Hung and Huo, 1985
		Kitthawee et al., 1999
		Pinto et al., 1993
		Pinto et al., 2003
Phosphogluconate dehydrogenase	Discrimination	Kimani-Njogu et al., 1998
		Kitthawee et al., 1999
		Unruh <i>et al.,</i> 1983
6-phosphoglucose dehydrogenase	Discrimination and detection	Castrovillo & Stock, 1981
Phosphoglucose isomerase	Discrimination	Atanassova et al., 1998
		Hung & Huo, 1985
		Unruh et al., 1983

Genomic sequences tend to repeat

Repetitive DNA sequences exist in several to millions of copies per cell (Hoy, 2003), which makes them useful for PCR targeting. For reasons that are not well understood, they tend to exhibit concerted evolution, i.e. non-independent evolution resulting in sequence similarity that is greater within than among species (Elder & Turner, 1995). They are therefore also useful for species differentiation.

Ribosomal RNA genes (rDNA) exist in arrays on one or more chromosomes and typically number in the thousands of copies per cell (Elder & Turner, 1995). The nuclear ribosomal DNA array in eukaryotes comprises three genes named on the basis of their sedimentation rates, 18S, 5.8S and 28S, preceded by an external transcribed spacer and separated by the ITS-1 and ITS-2 internal transcribed spacers; adjacent copies of the array are separated by a non-transcribed spacer (NTS) or intergenic spacer (Hillis & Dixon, 1991). 12S and 16S rDNA genes are found in the mitochondrial genome. Because different regions of the array evolve at different rates, they are useful for resolving differences at different taxonomic levels, with 16-18S genes being highly conserved and the 28S less so. The non-coding sequences also evolve, the NTS very rapidly, the nuclear ITS regions a little less so (Hillis & Dixon, 1991). One potential drawback is that, at least for ITS-2, there may be significant intraspecific variation in sequences (Allemand et al., 2002). Furthermore, intra-individual sequence variation may sometimes exceed inter-individual differences (Stouthamer et al., 1999; Alvarez & Hoy, 2002 and references cited therein), making discrimination of closely related populations problematical. The ITS-2 region discriminated some but not all of a suite of closely related Trichogramma (Hymenoptera: Trichogrammatidae) species (Stouthamer et al., 1999, 2000).

Mitochondrial sequences (mtDNA) are also found at high copy numbers per cell, typically hundreds or thousands (Hoy, 2003). A catalogue of conserved primers (Simon *et al.*, 1994) makes it a simple matter to 'fish out' specific mtDNA regions for sequencing to support specific primer design. Mitochondrial DNAs usually evolve three times faster than nuclear DNA because of inefficient repair mechanisms (however, ITS-1 may diverge at a more rapid rate than mtDNA; Elder & Turner, 1995). Since there is a finite risk that a putative amplified mtDNA sequence could actually be a nuclear 'pseudogene' evolving at a different rate than its mitochondrial ancestor, phylogenetic inferences and hypotheses about divergence times (cf. Chen *et al.*, 2002) employing mtDNA sequences must be made with care (Sunnucks & Hales, 1996; Benasson *et al.*, 2001).

Rapidly evolving tandem simple sequence repeats (SSRs) of 1-6 bp, sometimes referred to as microsatellites, are found throughout the coding and non-coding portions of genomes of all organisms. Their functions and mechanisms of evolution, and neutrality or selectivity, appear to be varied (Li et al., 2002); they also differ from other repeating sequences in being non-homologous, so that different copies within a given genome have different flanking sequences. Vanlerberghe-Masutti & Chavigny (1997) developed a species-specific microsatellite-based PCR assay for aphelinind parasitism; Hufbauer et al. (2001, 2004) used microsatellites to document reduced genetic variability in introduced populations of Aphidius ervi Haliday (Hymenoptera: Braconidae). Microsatellites of Diaretiella rapae (M'Intosh) (Hymenoptera: Aphidiidae) were used by Baer et al. (2004) to demonstrate lack of polyphagy, and by Loxdale & Macdonald (2004) to study movement of winged individuals at the farm scale.

In addition to microsatellites, the genome contains DNA comprising tandem arrays rich in A+T or C+G sequences, largely associated with chromosomal heterochromatin and centromeres (Elder & Turner, 1995; Hoy, 2003), and variously referred to as satellite or moderately repetitive DNA. Greenstone & Edwards (1998) used a 438 bp squashblot hybridization probe from a genomic library containing 53, 42, and 29 bp repeats to detect braconid endoparasitism. Similarly, Landais *et al.* (2000) used a genomic library to devise a 385 bp squash-blot hybridization probe containing 8–43 bp repeating motifs to differentiate and detect a suite of trichogrammatid parasitoids.

DNA assays

The simplest assays utilize complementary strands of the target sequences as hybridization probes. Such assays can detect small quantities of DNA if they employ probes comprising moderately repetitive sequences from the target species. Greenstone & Edwards (1998) developed such a probe that was able to detect early first instars, but not eggs, of the braconid *Microplitis croceipes* (Cresson) in lepidopteran host larvae in a squash-blot assay. Using the same approach, Landais *et al.* (2000) could detect the egg of *Trichogramma brassicae* (Bezdenko) 7 days post-parasitization within the host egg.

The most sensitive DNA assays employ PCR, which can amplify just a few copies of the target sequence. PCR using both random and specific primers has been applied to both parasitoid species discrimination and detection.

Random amplified polymorphic DNA markers (RAPDs)

RAPD markers are generally thought to sample highly polymorphic non-coding DNA (Black, 1993), to be phenotypically dominant or less often co-dominant, and to conform to Mendelian segregation patterns (Vaughn & Antolin, 1998); if co-dominant, i.e. having one strand with a large insertion or deletion, they can be detected using single-strand conformation polymorphism (SSCP) techniques (Sunnucks *et al.*, 2000). This permits population genetic analyses (but see Kazmer *et al.*, 1995, for caveats).

RAPDs were more successful than isozyme analysis in separating three *Muscidifurax* species (Hymenoptera: Pteromalidae) (Antolin *et al.*, 1996). RAPDs have effectively differentiated closely related congeneric species and strains within species (Baldanza *et al.*, 2001; Kirk *et al.*, 2000; Barnay *et al.*, 2001). Edwards & Hopper (1999) used RAPDs to differentiate hymenopterous parasitoids of differing genotypes emerging from individual host larvae, while Edwards & Hoy (1995) used them to distinguish pesticide-resistant from wild-type parasitoids of the same species. Edwards & Hoy (1993) looked for presence-absence rather than fragment-size polymorphisms, which greatly facilitates the scoring of differences between populations. Crossing studies showed that some but not all bands were inherited in Mendelian fashion as dominant traits.

RAPDs suffer from a reputation for lack of reproducibility both within one laboratory and from one laboratory to another (Black, 1993; Kazmer *et al.*, 1995; Taylor & Szalanski, 1999; Hoy *et al.*, 2000). Occasional contaminants, e.g. bacterial or protozoan symbionts, may produce bands that can be mistakenly identified as polymorphisms for the targets of interest (Black, 1993). There is also no reason to expect that similar-sized bands in different species are homologous (Black, 1993), a problem that does not arise with specific PCR approaches (below). RAPDs can be used as a discovery tool to find single-locus, species-specific sequences (see 'SCARS,' below under 'Specific PCR').

Black *et al.* (1992) provide the only report of latency in detection of parasitization by RAPD-PCR; parasitism in their aphid–parasitoid system could not be detected until six days post-parasitization.

Specific PCR

The availability of universal primer sequences for numerous genomic regions (Simon *et al.*, 1994) has facilitated the isolation and sequencing of DNA to enable design of parasitoid species-specific PCR primers. Table 2 summarizes the use of various genomic regions for species-specific differentiation and detection of parasitoid DNA. Not surprisingly, DNAs of hyperparasites as well as of primary parasitoids are detectable by PCR (Chen *et al.*, in press). Table 3 lists parasitoid species for which specific primers have been designed.

Although it is the newest technology to be applied to this field, specific PCR has been used more than any of the other approaches for parasitoid detection. Consequently, there is quite a bit of information on the sensitivity and time course of detectability of parasitization. Sensitivity is typically measured as wasp-equivalents of DNA detectable, generally in a reaction mixture containing host DNA (Zhu & Greenstone, 1999). Sensitivities on the order of 10^{-3} to 10^{-4} wasp equivalents are common (Zhu & Greenstone, 1999; Zhu *et al.*, 2000; Ratcliffe *et al.*, 2002), and sensitivities as low as 10^{-6} to 10^{-7} have been achieved (Zhu *et al.*, 2004b; Chen *et al.*, in press). This makes it possible, for example, to detect a single *Trichogramma* egg in the egg of its host (Li & Shen, 2002).

These high sensitivities also correspond to very short detection latencies, usually no more than 24 h post-parasitization (Zhu & Greenstone, 1999; Zhu *et al.*, 2000; Ratcliffe *et al.*, 2002; Zhu *et al.*, 2004b). Latencies less than 24 h have frequently been found (Amornsak *et al.*, 1998; Zhu & Williams, 2002; Persad *et al.*, 2004; Weathersbee *et al.*, 2004), and latencies longer than 24 h appear to be the exception (Zhu & Williams, 2002; Jones *et al.*, 2005).

Although specific PCR is very powerful, often enabling detection of just a single base-pair difference, it is not always possible to adjust PCR conditions to enable discrimination of all species of interest. In such cases, cutting the PCR product with a restriction enzyme may resolve these near-identical sequences. This process, known as restriction fragment length polymorphism (RFLP) analysis (also called restriction endonucease (REN) analysis; Escribano et al., 2000), requires knowledge of the target sequence so that restriction sites can be mapped. RFLP of a variety of coding and non-coding sequences has been widely used to differentiate closely related parasitoid species (Sappal et al., 1995; Silva et al., 1999; Taylor & Szalanski, 1999; Stouthamer et al., 1999; Tilmon et al., 2000; Allemand et al., 2002; Asfaq et al., 2004; Borghuis et al., 2004). Prinsloo et al. (2002) used RFLP and two genomic regions, ITS-2 and 16s rDNA, to distinguish three sibling species of Aphelinus (Hymenoptera: Aphelinidae).

In 1993, Paran & Michelmore published a technique, which they dubbed sequence characterized amplified regions (SCARS), in which PCR products are excised from a RAPD gel, sequenced, and used for primer design. This approach was used by Zhu *et al.* (2004a) to distinguish pesticide-resistant and -susceptible strains of a pteromalid parasitoid by isolating and cloning strain-specific fragments amplified by RAPD–PCR, and then making PCR primers to amplify these specific but theretofore unknown fragments.

Recommendations

This review has identified four distinct molecular approaches that have been applied to the discrimination and detection of insect parasitoids, and that may be appropriate for assessing the extent of insect parasitism. Immunoassay, especially with monoclonal antibodies, is sensitive and has great powers of discrimination, enabling stage- as well as species-specificity. Nevertheless, the relative paucity of studies employing it, especially given the

Region	Application	References
Cytochrome B	Discrimination	Daza-Bustamente et al., 2002
Cytochrome oxidase I	Discrimination	Baer <i>et al.</i> , 2004
		Hufbauer et al., 2004
	Discrimination and detection	Ashfaq et al., 2004
		Agustí <i>et al.</i> , 2005
		Persad <i>et al.</i> , 2004
		Tilmon <i>et al.</i> , 2000
Cytochrome oxidase II	Discrimination	Baer <i>et al.</i> , 2004
		Borghuis <i>et al.</i> , 2004
		Hufbauer <i>et al.</i> , 2004
ITS-1	Discrimination	Chang <i>et al.</i> , 2001
		Hufbauer <i>et al.</i> , 2004
		Orrego & Agudelo-Silva, 1993 Sappal <i>et al.</i> , 1995
		11
	Discrimination and detection	Taylor & Szalansi, 1999
	Discrimination and detection	Gariepy <i>et al.</i> , 2005 Ratcliffe <i>et al.</i> , 2002
Cytochrome oxidase I/II	Discrimination	
cytochrome oxidase 1/11	Discrimination	Daza-Bustamente <i>et al.</i> , 2002
ITS-2	Discrimination	Schneider <i>et al.,</i> 2003 Allemand <i>et al.,</i> 2002
115-2	Discrimination	Alvarez & Hoy, 2002
		Campbell <i>et al.</i> , 1993
		Hufbauer <i>et al.</i> , 2004
		Persad <i>et al.</i> , 2004
		Sappal <i>et al.</i> , 1995
		Silva <i>et al.</i> , 1999
		Stouthamer <i>et al.</i> , 1999
		Stouthamer <i>et al.</i> , 2000
	Discrimination and detection	Amornsak <i>et al.</i> , 1998
		Ashfaq et al., 2004
		Erlandson <i>et al.</i> , 2003
		Gariepy <i>et al.</i> , 2005
		Li & Shen, 2002
		Prinsloo et al., 2002
		Zhu & Greenstone, 1999
		Zhu & Williams, 2002
		Zhu et al., 2000
		Zhu et al., 2004b
28s rDNA	Discrimination	Campbell et al., 1993
		Persad et al., 2004
		Sappal <i>et al.</i> , 1995
18s rDNA	Discrimination and detection	Suckling et al., 2001
		Weathersbee et al., 2004
16s rDNA	Discrimination	Baer et al., 2004
		Chen <i>et al.</i> , 2002
		Persad et al., 2004
	Discrimination and detection	Chen et al., in press
		Jones <i>et al.</i> , 2005
		Prinsloo et al., 2002
12s rDNA	Discrimination	Persad et al., 2004
NADH	Discrimination	Persad et al., 2004
Actin genes	Discrimination	Hoy et al., 2000
Esterase-like enzyme	Discrimination	Zhu et al., 1999
Satellites	Discrimination and detection	Greenstone & Edwards, 1998
Microsatellites		Landais et al., 2000
	Discrimination	Amornsak et al., 1998
		Hufbauer et al., 2001
		Jensen et al., 2002
		Loxdale & MacDonald, 2004
		MacDonald et al., 2003
Unknown sequences	Discrimination	Vanlerbeghe & Masutti, 1997 Zhu <i>et al.</i> , 2004a

Table 2. DNA regions used in specific PCR and PCR-RFLP.

maturity and broad application of the technology to other disciplines, suggests that it may not be practical for biocontrol practitioners. RAPDs have been used for many applications, but only once (Black *et al.*, 1992) for detection, and exhibited low sensitivity in that instance. Given its poor laboratory-to-laboratory reproducibility, it is most useful for

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Species	Family	Region	References
Ageniapsis citricola	Encyrtidae	Actin genes	Hoy et al., 2000
Alloxysta xanthopis	Charipidae	16s rDNA	Chen et al., in press
Anaphes iole	Mymaridae	ITS-2	Zhu & Williams, 2002
Anisopteromalus calandrae	Pteromalidae	Esterase-like enzyme	Zhu et al., 1999
		Uncharacterized	Zhu <i>et al.</i> , 2004a
Aphelinus abdominalis	Aphelinidae	Microsatellite	Vanlerberghe-Masutti & Chavigny, 1997
A. albipodus		ITS-2	Zhu & Greenstone, 1999
A. asychis			
A. gossypii		18s rDNA	Weathersbee <i>et al.</i> , 2004
A. hordei		ITS-2	Zhu <i>et al.</i> , 2000
		ITS-2, 16s rDNA	Prinsloo <i>et al.</i> , 2002
A. varipes		ITS-2	Zhu & Greenstone, 1999
Aphidius colemani	Aphidiidae	ITS-2	Zhu <i>et al.,</i> 2000
A. ervi		CO I-II, CO B	Daza-Bustamante <i>et al.</i> , 2002
	D 11	Microsatellite	Hufbauer <i>et al.</i> , 2001
Cotesia congregata	Braconidae	Microsatellite	Jensen <i>et al.</i> , 2002
Diaretiella rapae		Microsatellite	Loxdale & Macdonald, 2004
Dendrocerus carpenteri	Megaspilidae	16s rDNA	Chen <i>et al.</i> , in press
Dolichogenidia tasmanica	Braconidae	18s rDNA	Suckling <i>et al.</i> , 2001
Leiophron argentinensis		ITS-2	Zhu <i>et al.,</i> 2004a
L. uniformis			D 1 (1 0001
Lipolexis oregmae	Aphidiidae	ITS-2	Persad et al., 2004
L. scutellaris		18s rDNA	Weathersbee <i>et al.</i> , 2004
Lydella thompsoni	Tachinidae	COI	Agustí et al., 2005
Lysiphlebus testaceipes	Aphidiidae	ITS-2	Persad et al., 2004
		18s rDNA	Weathersbee <i>et al.</i> , 2004
14 116 4	D: 1:1	16s rDNA	Jones <i>et al.</i> , 2005
Muscidifurax raptor	Pteromalidae	ITS-1	Ratcliffe <i>et al.</i> , 2002
M. raptorellus			
M. zaraptor	Dr. 1:1		
Nasonia vitripennis	Pteromalidae	ITS-1	Ratcliffe <i>et al.</i> , 2002
Peristenus digoneutis	Pteromalidae	ITS-1, ITS-2	Erlandson <i>et al.</i> , 2003
P. howardi			Gariepy <i>et al.</i> , 2005
P. pallipes			Zhu et al., 2004b
P. pseudopallipes			Erlandson <i>et al.</i> , 2003
			Gariepy <i>et al.</i> , 2005
Detusious			Zhu et al., 2004b
P. stygicus			Erlandson <i>et al.</i> , 2003
			Gariepy et al., 2005 Zhu et al. 2004b
Pseudoperichaeta nigrolineata	Tachinidae	CO I	Zhu <i>et al.,</i> 2004b Agustí <i>et al.,</i> 2005
Spalangia calcitrans	Pteromalidae	ITS-1	Ratcliffe <i>et al.</i> , 2002
S. cameroni	Tteromandae	113-1	Ratellife et ul., 2002
S. endius			
S. nigroaena			
Trichomalopsis sarcophagae	Pteromalidae	ITS-1	Ratcliffe et al., 2002
Urolepis rufipes	Pteromalidae	ITS-1	Ratcliffe <i>et al.</i> , 2002
Trichogramma australicum	Trichogrammatidae	ITS-2	Amornsak <i>et al.</i> , 1998
T. brassicae	menogrammatidae	Satellite	Landais <i>et al.</i> , 2000
1. 01400teac		ITS-1, ITS-2	Sappal <i>et al.</i> , 1995
T. chilonis		ITS-1, 113-2 ITS-1	Chang <i>et al.</i> , 2001
T. deion		ITS-2	Stouthamer <i>et al.</i> , 1999
T. dendrolimi		ITS-2	Li & Shen, 2002
T. minutum		ITS-1, ITS-2	Sappal <i>et al.</i> , 1995
T. ostriniae		ITS-1	Chang <i>et al.</i> , 2001
			0 ,
		ITS-2	Stouthamer et al., 1999
T. pretiosum		ITS-2 ITS-2	Stouthamer <i>et al.,</i> 1999 Li & Shen, 2002

a particular short-term project, or for long-term studies of a particular system and run by a single laboratory.

Allozyme electrophoresis and specific PCR, on the other hand, exhibit excellent reproducibility, and, unlike RAPDs, use markers whose homologies across species can be verified by activity and sequence, respectively. The compilation in table 1 suggests that perhaps a half dozen enzyme classes, led by esterases, phosphoglucomutase, and malate dehydrogenase, are adequate for separating most parasitoid taxa by allozyme electrophoresis. When faced with having to make a choice of which enzyme system to use to assay a large number of field-collected hosts, Walton *et al.* (1990b) chose the esterase system as the 'most versatile' system for parasitoid detection in their system. The few studies of sensitivity, all involving aphid parasitoids, have given detection times greater than 3 days post-parasitization. As indicated by Richardson *et al.* (1986), many factors influence the appearance and interpretation of banding patterns. Furthermore, particular host-parasitoid systems may have their own quirks, requiring a certain amount of experience for correct interpretation (H.D. Loxdale, personal communication).

As table 2 shows, much of the genome has been used for discrimination and/or detection of parasitoids by specific PCR. Despite some concerns about intra-individual variation, ITS-2 has been most used for detection, followed by cytochrome oxidase. Cytochrome oxidase I (COI) has been proposed as a universal barcode for all animals, and there is general agreement that COI will achieve this for most animal species (see http://www.barcodinglife.com). Therefore, broader employment of COI would increase the utility of markers developed for biocontrol use. In cases where sequence similarity is so high, for example in some groups of parasitoid sibling species, that COI cannot resolve differences (e.g. see Alvarez & Hoy, 2002), RFLP of COI targets, or amplification of other genomic sequences, may then be necessary.

Specific PCR has exhibited extraordinary sensitivity, detecting as little as 10^{-7} wasp equivalents, enabling detection of parasitoid eggs or larvae in their hosts within 24-48 h of oviposition in all studied cases, hence an advance over allozymes analysis. The fact that many pests, and in some cases their parasitoids, are nearly cosmopolitan, means that primer sequences once developed in any laboratory may be used by other biocontrol practitioners in their field situations. Finally, the maturity of the PCR technique, and the very large pool of talented entomologists able to develop, employ, and trouble-shoot this methodology, augurs well for PCR to become the dominant assay method in the near term. Optimization of multiplex PCR protocols (Gariepy et al., 2005), also utilized in predator-prey research (e.g. Harper et al., 2005), will simplify application of the methodology to multi-parasitoid systems.

Even so, DNA technologies are very rapidly evolving, and we may expect before this first decade of the new millennium is out to see DNA hybridization assays employing nanotechnology approaches exhibiting very high sensitivities and specificities (e.g. see Taton *et al.*, 2000 and Parks *et al.*, 2002). When these approaches are standardized and applied to parasitism research, we will not have to run PCR assays to assess parasitism rates: we shall use our thermocyclers only to fish out species-specific primers, which we shall in turn use to design species-specific probes for use in sensitive, 'quick-and-dirty' hybridization assays.

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