INVITED REVIEW

Plant Architecture: A Dynamic, Multilevel and Comprehensive Approach to Plant Form, Structure and Ontogeny

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• Background and Aims The architecture of a plant depends on the nature and relative arrangement of each of its parts; it is, at any given time, the expression of an equilibrium between endogenous growth processes and exogenous constraints exerted by the environment. The aim of architectural analysis is, by means of observation and sometimes experimentation, to identify and understand these endogenous processes and to separate them from the plasticity of their expression resulting from external influences.

• Scope Using the identification of several morphological criteria and considering the plant as a whole, from germination to death, architectural analysis is essentially a detailed, multilevel, comprehensive and dynamic approach to plant development. Despite their recent origin, architectural concepts and analysis methods provide a powerful tool for studying plant form and ontogeny. Completed by precise morphological observations and appropriated quantitative methods of analysis, recent researches in this field have greatly increased our understanding of plant structure and development and have led to the establishment of a real conceptual and methodological framework for plant form and structure analysis and representation. This paper is a summarized update of current knowledge on plant architecture and morphology; its implication and possible role in various aspects of modern plant biology is also discussed.

Key words: Plant morphology, plant architecture, level of organization, growth, branching, differentiation, morphogenetic gradients, physiological age, meristem, annual shoot, phenotypic plasticity, ontogeny, phase change.

INTRODUCTION

Progress in our understanding of plants has increased dramatically in recent decades and research in this domain has given rise to analytical, methodological and theoretical innovations on various aspects of plant science (Sattler and Rutishauser, 1997; Hedden et al., 2002; Turnbull, 2005).

Plant form, development and evolution have been analysed under the functional view of biomechanics (Niklas, 1992, 2005; Diggle, 2003; Friedmann and Harder, 2004, 2005). The importance of phenotypic plasticity, phenology and crown architecture in evolutionary plant ecology or in the understanding of community and stand structure, functioning and production has been stressed by several authors (Givnish, 1984; King, 1998; Diggle, 1999, 2002; Huber et al., 1999; Alpert and Simms, 2002; Novoplansky, 2002; Wright et al., 2002; Oborny, 2004; Sachs, 2004; Damascos et al., 2005; de Kroon et al., 2005; Pearcy et al., 2005; Wolfe and Mazer, 2005).

Since its introduction and definition by von Goethe (1790), plant morphology has had a successful history and it is commonly accepted that plants are modular organisms that develop by the repetition of elementary botanical entities whose morphological, dimensional, functional and anatomical features change during ontogeny and according to several processes variously called heteroblasty, phase change, life stages, maturation, ageing, age states or morphogenetic progression (Goebel, 1900; Troll and Rauh, 1950; Robbins, 1957; Wareing, 1959, 1961; Nozeran et al., 1971, 1982; Nozeran, 1978, 1984; Gatsuk et al., 1980; Greenwood, 1987, 1995; Poethig, 1990; Jones, 1999, 2001; Claßen-Bockhoff, 2001; Kaplan, 2001). As plant morphology deals with plant form and/or structure and with their temporal and/or topological changes during ontogeny and even phylogeny, it is therefore relevant to practically all the disciplines of modern plant biology cited above (Sattler, 1978; Roloff, 1988; Sattler and Rutishauser, 1997; Gleichner, 1998; Claßen-Bockhoff, 2000, 2005; Kaplan 2001; Scotland et al., 2003; Wiens, 2004; Mueller, 2006).

The study of plant architecture emerged as a new scientific discipline some 30 years ago, and derived, in several ways, from earlier works on plant morphology (Halle and Oldeman, 1970; Oldeman, 1974; Halle et al., 1978). An original feature of architectural studies is that they were initiated in tropical regions and were, at first, concerned with the analysis of the aerial vegetative structure of tropical trees (Halle and Oldeman, 1970). Since their definition, architectural concepts have provided powerful tools for studying plant form or even tropical forest structure and the understanding of its dynamics (Oldeman, 1974, 1983, 1990; Halle et al., 1978; Vester, 1997). Investigations based on these concepts quickly spread to temperate species (Edelin, 1981; Caraglio and Edelin,
Growth process

The primary growth of a plant is the result of several processes that can be grouped into two distinct, but co-ordinated morphogenetic events: organogenesis and extension (Champagnat et al., 1986). The inception of new organs (organogenesis) results from the functioning of undifferentiated cells constituting the apical meristem (Fig. 1A). Located at the tip of a stem, this meristem forms, when in an active phase, small cell masses with

![Diagram of plant growth processes](http://aob.oxfordjournals.org/)

**Fig. 1.** Shoot apex (A) and stem (B) organization. Each leafy axis (B) ends in an apical meristem frequently protected in an apical bud (A). Each stem comprises a succession of metamers, i.e. the set composed by (1) one internode, (2) the node (i.e. insertion point of the leaves on a stem) located at its tip and (3) the corresponding one or several leaves and associated lateral buds (in grey on A; White, 1979; Caraglio and Barthélémy, 1997).
different potentialities (Lyndon, 1988; Nougare`de, 2001); these masses develop into embryonic leaves, then leaves on elongated stems. The insertion zone of a leaf on the stem is referred to as a node and the stem portion which separates two successive nodes is called the internode. According to species and phyllotaxis, one, two or more leaves may be inserted at one node and one to several, named supernumerary buds (see Fig. 8) may develop at the axil of each individual leaf. A stem can thus be considered as a succession of internodes and the entity formed by a node, associated with its leaf (or leaves) and axillary bud(s) plus the subtending internode, represents the basic structural unit of the plant body commonly called the metamer or phytomer (White, 1979; Barlow, 1989). During growth, the superposition and repetition of this elementary entity builds up the leafy axis (Fig. 1B).

Plant growth may be considered in several ways according to the kind of organ and/or level of organization concerned (growth of a leaf, a stem, a fruit, the whole plant, etc.). As we are dealing here with the topological edification of the stem and macroscopic aspects of plant growth, we will focus mainly on the extension process of the leafy axis or shoot and will neither detail organogenesis processes nor consider secondary (sensu Fahn, 1967) or radial growth.

Determinate vs. indeterminate growth (Fig. 2). In many plant species, the apex may abscise or abort after some period of functioning (Garrison and Wetmore, 1961; Millington, 1963; Puntieri et al., 1998, 1999) or it may transform into a specialized structure (flower, inflorescence, spine, tendril, parenchymatous cells, etc.) lacking further extension capacity. In these cases, the axis is considered to have a determinate growth (Fig. 2A–C; Hallé et al., 1978; ‘definite extension’, Bell, 1991). By contrast, indeterminate growth (Hallé et al., 1978) or indefinite extension (Bell, 1991) refers to an axis on which apical meristem indefinitely maintains its growth potential (Fig. 2D). As the indefinite functioning of an apex is always limited at least by the limited life span of the plant it belongs to, this ultimate term is somewhat ‘theoretical’ and a misuse of the language (Guinochet, 1965). Nonetheless, this notion is useful and justified by the necessity to describe and name this phenomenon.

Rhythmic vs. continuous growth. Hallé et al. (1978) distinguished shoots which have no marked endogenous cessation of extension (continuous growth) from shoots which have marked endogenous periodicity and cessation of extension (rhythmic growth; Hallé and Martin, 1968). Although organogenesis may be considered or known, these two growth patterns are generally more concerned with extension.

Continuous extension (Fig. 3A, B) is quite a rare phenomenon in the field and has generally been observed and described mostly in uniform equatorial climates or environments (mainly palm or mangrove trees; Venkatanarayana, 1957; Rees, 1964; Gill and Tomlinson, 1971) or for some herbs native to temperate (Bell, 1991) or tropical (Blanc, 2002) regions. In all these cases, plants exhibit a more or less constant production of leaves and/or shoots throughout the year. For the mangrove trees *Rhizophora mangle* (Gill and Tomlinson, 1971), growing in the subtropical climate of Florida, it was shown that the extension of each axis may have a fluctuating rate according to the seasonal climatic conditions, without ever being interrupted completely (Fig. 3B). In some cases, the endogenous nature of continuous extension may be ‘masked’ by fluctuating environmental conditions in the field and can be revealed experimentally by growing plants in favourable and constant conditions, as was demonstrated for several temperate Cupressaceae from the Southern Hemisphere (Grosfeld and Barthélemy, 2004). It has been shown that plants with continuous extension may also present a continuous production of new embryonic leaves, i.e. organogenesis (Gill and Tomlinson, 1971). As far as it has been documented, when extension is continuous the resulting stem is generally quite homogeneous and successive metamers and their constitutive elements are more or less of the same type and size along the elongated axis (Fig. 3A, B, right; Tomlinson and Gill, 1973; Hallé et al., 1978).
Rhythmic extension of leafy axes is a far more frequent growth pattern in plants, regardless of their geographical origin, and is expressed by an alternation of periods of rest in meristem activity and periods of active shoot extension or ‘growth flushes’ (Fig. 3C). This rhythmic growth has been thoroughly studied by numerous authors (Alvim, 1964; Kozlowski, 1971; Hallé et al., 1978; Puntieri et al., 1998; Sabatier and Barthélémy, 1999). Hallé and Martin (1968), in their detailed study of the tropical tree *Hevea brasiliensis*, defined the ‘growth unit’ as the portion of an axis which develops during an uninterrupted period of extension. A growth unit is generally easy to identify on the elongated stem as the limit between two growth units is usually marked by a zone of short internodes and/or cataphylls (i.e. scale leaves) corresponding to the protective organs of the bud from which it derives (Fig. 4). Even if very common in rhythmically growing axes, this alternation of cataphylls and leaves (termed articulate growth by Tomlinson and Gill, 1973; Fig. 3C right, Fig. 4) may not be obvious in some species, and other morphological or macro-anatomical markers may sometimes be used in addition, for the *a posteriori* identification of a rhythmically elongating axis (Fig. 5).

As with continuous growth, the identification of the endogenous nature of rhythmic growth, for plants naturally growing in seasonal climates, requires experimental verification in controlled, constant and favourable environmental conditions (Greathouse et al., 1971; Lavarenne, 1971; Payan, 1982; Parisot, 1985). Regardless, the rhythmic (or continuous) pattern of a shoot or axis growth must
be checked by periodic measurement of its length. In practice, most often only the extension component of growth is known, such that some authors have proposed the use of the term ‘unit of extension’ (Gill and Tomlinson, 1971; Halle` et al., 1978) rather than growth unit, even though the two terms are synonymous. When organogenesis is known, it has been shown that the rhythmic extension of the stem may be combined either with a continuous (Bond, 1942) or more frequently, a rhythmic (Halle` and Martin, 1968; Gill, 1971; Puntieri et al., 2002a; Sabatier et al., 2003b) organogenesis pattern. In the latter case, it has been proposed that the ‘unit of morphogenesis’ is the axis portion corresponding to an uninterrupted phase of organogenesis (Halle` and Martin, 1968). Despite its relevance for the understanding of growth patterns in plants, the nature of the unit of morphogenesis is unknown in most cases partly owing to the long and tedious experimental work needed for its identification.

Preformation and neoformation. In the case of rhythmic growth, all the metamers and organs of the future elongated shoot may be present at an embryonic stage in a bud before the elongation of the shoot deriving from it; in this case the shoot is referred to as ‘preformed’ and its constitutive organs as ‘preformed organs’ or ‘preformation’ (Halle` et al., 1978; ‘early leaves’, Critchfield, 1960; ‘fixed growth’, Lanner, 1976). The duration of preformed organs at an embryonic stage in a bud may vary from several days or weeks (Halle` and Martin, 1968; Sabatier et al., 1995) to several years (Aydelotte and Diggle, 1997; Diggle, 1997; Meloche and Diggle, 2001). In other cases, more organs than those included at an embryonic stage in the bud are elongated: these supplementary, non-preformed elements are referred to as ‘neoformed organs’ (i.e. ‘neoformation’ sensu Halle` et al., 1978, or ‘late leaves’, Critchfield, 1960; ‘free growth’, Jablanczy, 1971; Lanner, 1976). As a consequence stems or shoots may comprise only preformed metamers (Critchfield, 1960; Rivals, 1965, 1966; Gill, 1971; Roloff, 1985; Sabatier et al., 1995; Nicolini, 1998; Puntieri et al., 2000, 2002a, b; Souza et al., 2000) or, more rarely, may be entirely neoformed (Borchert, 1969; El-Morsy, 1991). In many cases, a preformed part can be followed by a neoformed part and thus give rise to a mixed shoot (Critchfield, 1960; Halle` and Martin, 1968; Kozlowski, 1971; Nitta and Ohsawa, 1998; Puntieri et al.,...
Annual shoot (Fig. 6). In rubber trees growing in equatorial conditions (Hallé and Martin, 1968), an axis with indeterminate growth forms a new growth unit about every 45 d and these successive units are morphologically identical (cf. Fig. 3C) so that the growth unit level itself is pertinent to describe the infrastructure of an axis and its morphological heterogeneity. In other tropical species, however, the timing of rhythmic extension during a whole calendar year is more complex. In *Ryania speciosa var. subuliflora* (Comte, 1993), for instance, the annual growth pattern corresponds to the extension of two growth units in a relatively short time, followed by a long resting phase, which again is followed by the rapid emission of a succession of two growth units. In some temperate species, shoot extension may occur in one, two or more successive events in a same growing season giving rise to an ‘annual shoot’ made up of one or a succession of several growth units or growth cycles (respectively monocyclism vs. polycyclism, i.e. Bugnon and Bugnon, 1951 or Lanner 1976, and see Caraglio and Barthélémy, 1997, for a critical and historical revision of these terms) occurring in a vegetative cycle (period between two marked seasons or two winters, Lanner, 1976). When several successive growth units are formed in the same annual vegetative cycle these growth units most often present distinctive features (Fig. 6); spring shoots and summer or additional shoots are frequently distinguished (Späth, 1912; Kozlowski, 1971; Cannell et al., 1976). In this situation, successive growth units produced in a same year are thus not identical and it is useful and pertinent to distinguish an annual-shoot level of organization.

According to the number of constitutive growth units of an annual shoot and according to the indeterminate vs. determinate nature and to the relative extent of preformation and neoformation in each of the constitutive growth units, several combinations may exist for different species or even for a single species, depending on external and/or internal conditions (Lanner, 1971, 1976; de Reffye et al., 1991; Costes, 1993; Fontaine et al., 1999; Heuret et al., 2000, 2003, 2006; Puntieri et al., 2000; Isik et al., 2001).

**Branching process**

Although the architecture of some vascular plants consists only of a single vegetative axis during their whole life span, most display a more complex architecture consisting of several axes, one derived from another by a repetitive process known as branching.

**Terminal vs. lateral branching.** An apical meristem (McManus and Veit, 2002) or an initial apical cell (Gifford, 1983) can directly split and give rise to two or more new sibling axes, which results in dichotomy or polytomy, respectively (Emberger, 1960). Frequently encountered in ferns and mosses (Emberger, 1960), this terminal branching (Gatin, 1924) is rarely expressed in angiosperms (see for monocotyledons: Schoute, 1909; Tomlinson, 1971; Fisher, 1976; Tomlinson and Poslusny, 1977; and for dicotyledons: Nolan, 1969; Boke, 1976; Iwamoto et al., 2005). It is important to mention that this phenomenon is, as far as we know, not encountered within gymnosperms.

In other cases, the branching process relies on the delimitation of a zone of embryonic cells just aside the initiated leaf, i.e. the axillary meristem. This lateral branching process is the most common one among angiosperms and gymnosperms. The resulting lateral branch is characterized by the presence of one or two (in respectively monocotyledons or dicotyledons) prophylls: the first foliar organs of the lateral axis (see Fig. 7A, B for their particular location).

In some plants, there are, at a single leaf axil, more than one axillary meristem, which are then called supernumerary or accessory buds (Sandt, 1925; Troll, 1937; Espagnac and Neuville, 1969; Altman and Goren, 1978; Bell, 1991; Fig. 8A, B). In this case, each individual lateral meristem may be identified by the position of its prophylls, which
also allows the distinction between this situation and the development of condensed and more complex lateral branching systems (i.e. complexes of secondary buds, Hallé et al., 1978), where second-order lateral buds may be present in the axils of main lateral bud’s prophylls (Fig. 8C).

Immediate vs. delayed branching (Fig. 9). Once initiated, an axillary meristem may remain dormant or can develop into a lateral axis. A lateral axis may elongate immediately after lateral meristem initiation or after a phase during which the lateral meristem remains inactive and very often protected in a lateral bud. These branching patterns are referred to, respectively, as immediate (i.e. ‘syleptic’ sensu Hallé et al., 1978; Müller-Doblies and Weberling, 1984; Wu and Hinkley, 2001) or delayed (‘proleptic’ sensu Hallé et al., 1978; ‘prolepsis’, Bell, 1991) branching. The terms immediate vs delayed branching should be preferred to the more traditional ‘syleptic’ vs. ‘proleptic’ branching (sensu Hallé et al., 1978) because of etymological and historical reasons, which have rendered the latter two terms ambiguous (see Caraglio and Barthélémy, 1997, for a critical review).

Morphologically, these two branching patterns may generally be identified a posteriori by the observation of the base of the lateral axis. Because of the immediate extension of lateral organs, branches with immediate development generally lack proximal cataphylls and present a relatively long most proximal internode termed a hypopodium (Fig. 9A; Tomlinson and Gill, 1973). Irrespective of the delay length, delayed branches present very short internodes and one or several cataphylls in their proximal portion, close to the point of insertion (Fig. 9B; Hallé et al., 1978). Thorough observation of the proximal part of lateral axes and a periodic observation and measurement of lateral meristem or bud size normally leaves no doubt as to the immediate or delayed nature of branching. In some cases, however, because lateral resting buds may be ‘hidden’, during the resting phase, in the axils of persistent basal foliar organs of the terminal bud (Guédès, 1975, 1980) or because lateral axes in an embryonic stage may already exist inside a resting bud (Champagnat, 1965; Roloff, 1985), the interpretation may be complicated or may lead to a misuse of the terminology (Caraglio and Barthélémy, 1997). Finally, in delayed branching it has been shown that the duration of the delay may be of several weeks to a year (Sabatier et al., 1995, 1998, 2003b; Puntieri et al., 1998; Heuret et al., 2003) and even several years (Fink, 1983; Nicolini et al., 2001).

Monopodial vs. sympodial branching. Depending on the indeterminate or determinate growth pattern of an axis, its branching pattern may respectively be monopodial (Emberger, 1960; or ‘monopodic’, Sachs, 1874), or sympodial (Emberger, 1960; or ‘sympodic’, Sachs, 1874). In the latter case, one, two or more branches may develop after the death, abscission, abortion or transformation of the apex, and the resulting sympodial branching pattern be qualified respectively as mono-, di- or polychasial. In plant architecture, the concept of module (‘article’ in

Fig. 7. The leaf, or leaves, of the first (proximal) node of a lateral shoot (A) are referred to as prophylls. In dicotyledons prophyll α and prophyll β are mainly in opposite and lateral position with respect to the plan formed by the axillary leaf (L) and the parent axis (P) (Salvia guaranitica, A). In monocotyledons, the first leaf (prophyll α) is often bicarinate and shows a particular arrangement (unidentified Poaceae, B): it is located in adaxial position between the lateral shoot (A) and its parent axis (P).

Fig. 8. Vertical succession of supernumerary (or accessory) buds in Juglans regia (A) and in Forsythia vulgaris (B). The arrangement of the prophylls (diagrams) distinguishes supernumerary buds from reduced branching systems as illustrated in Zelkova serrata (C). L, axillary leaf; Ls, leaf scar; P, parent axis; prophylls α and β or their scars, αs and βs, after abscission.
French) was defined for the first time by Prévost (1967) in her study of tropical Apocynaceae and then applied to ‘a leafy axis in which the entire sequence of differentiation is carried out from the initiation of the meristem that builds up the axis to the sexual differentiation of its apex’ (Halle, 1986). Because of the various causes of determinate growth (see above), this definition is, however, restrictive and the term ‘module’ increasingly refers to the portion of an axis edified by a single terminal meristem, which corresponds also to a ‘sympodial unit’ as used by Bell (1991).

Involving one or more lateral axes, a sympodial branching system is very often three dimensional. In some cases, however, a sympode may imply a linear succession of ‘modules’ or ‘sympodial units’ forming a so-called ‘pseudomonopodium’ as termed by German morphologists (Troll, 1937; Rauh, 1939). The resulting rectilinear structure mimics an axis edified by a single meristem with indeterminate growth and, if reproduced on all axes, may give rise to a totally sympodial branching system that resembles a monopodial one (Caraglio and Edelin, 1990; Bell, 1991). As a consequence, a rectilinear stem may thus be composed of a succession of metamers or growth units or annual shoots all produced by a single meristem or by a linear succession of sympodial units or modules. In a broader sense and following Room et al. (1994), the term ‘leafy axis’ or ‘axis’ will identify not only a structure edified by a single meristem as initially considered by Hallé et al. (1978) but also a rectilinear stem, whatever its intrinsic mode of construction.

Branched system and branching order (Fig. 10). The description of a branched system implies the use of a precise topological terminology. In plant architecture analysis it is usual to use ordinal numbers and to consider the main stem arising from seed as the order 1 axis (Fig. 10A; Hallé et al., 1978; Barthélémy et al., 1989, 1991) whereas the axes it gives rise to are referred to
order 2 axes and so on. In a sympodial system, a rigorous use of this terminology will lead to the reference of the successive sympodial units as axis orders 1, 2, 3, etc. In order to be coherent with our broad axis definition (see above), each rectilinear succession of modules, even though not strictly edified by a single meristem, will be considered as an axis and will represent an ‘apparent branching order’ (Fig. 10A, C).

**Rhythmic vs. continuous or diffuse branching.** Defined and used for the first time for plant architecture description, these terms contribute to the characterization and definition of architectural models (Halle´ and Oldeman, 1970; Halle´ et al., 1978) and take into account the topological distribution of sibling axes on a parent axis. Depending on whether all the axillary meristems of a stem develop into lateral axes, or whether lateral axes are grouped as distinct tiers with an obvious regular alternation of a succession of unbranched and branched nodes on the parent stem, branching is respectively referred to as continuous or rhythmic. In some cases, neither all nodes of a parent axis are associated with a lateral axis nor is there an obvious regular distribution of branches in tiers, and the branching pattern is then called ‘diffuse’. As revealed in Cupressaceae by qualitative observations (Courtot and Baillaud, 1961; Baillaud, 1999) and, in recent years by sophisticated mathematical methods (Guédon et al., 2001; Grosfeld, 2002; Heuret et al., 2002), a diffuse branching pattern may not mean an unorganized distribution of sibling shoots on a parent shoot but may indicate a predictable, precise and subtle branching organization.

**Acrotonic vs. mesotonic or basitonic branching (Fig. 11).** In order to describe the positional preferential development of lateral axes on a vertical parent axis or shoot, Troll (1937) and Rauh (1939) distinguished three modalities that were grouped under the German expression ‘longitudinale Symmetrie’. Acrotony (Fig. 11A, B) is the prevalent development of lateral axes in the distal part of a parent axis or shoot, and depending on whether branching is monopodial or sympodial, Rauh (1939) and Champagnat (1947) termed it, respectively, ‘primary’ or ‘secondary’ acrotony. In the initial definition of acrotony, the parent axis was always longer than the sibling ones, but Bell (1991) used the term ‘acrotonic branching’ for the distal position of the largest lateral branch, independent of its size relative to parent stem. Basitony (Fig. 12) was at first (Troll, 1937) considered as the preferential development of lateral axes in the basal part of a vertical stem. Bell (1991) used the term ‘basitonic branching’ when the proximal branches grow larger than the distal ones. Finally, mesotony (Fig. 11C, D) is used for a privileged development of branches in the median part of a shoot or axis.

Clearly included in the definition (Bell, 1991) or implicitly shown in the illustrations (Troll, 1937; Rauh, 1939), the topological lateral arrangement of branches along the parent axis is often associated with an increasing or decreasing gradient in length and/or vigour of the branches. As discussed by Caraglio and Barthélémy (1997), these two points must nevertheless be considered separately as the diversity of their expression in plants shows that there is neither automatic nor direct correlation between the privileged position and the relative vigour of lateral branches (compare Fig. 11A with Fig. 11B, or Fig. 11C with Fig. 11D). As all kinds of combinations may be found in the plant kingdom, we would prefer the terms acrotony, basitony and mesotony to be used only in reference to the privileged localization of branches on a
parent shoot (respectively distal part, proximal part and median position) without reference to their relative vigour or length, which can be given in precise terms in addition.

Acrotony or basitony are frequently considered as two fundamental phenomena underlying, respectively, the ‘arborescent’ or ‘bushy’ growth habit (Troll, 1937; Rauh, 1939; Champagnat, 1947; Barnola and Crabbe, 1991). Nevertheless these authors refer mainly to the acrotropic branching of growth units or annual shoots in the arborescent case whereas they consider the proximal branching at the base of the whole individual when considering bushy plants. As discussed by Caraglio and Barthelémy (1997) either acrotropic or basitropic branching may characterize the growth units or shoots of either tree or bush, whereas basal sprouts may also be an adaptive strategy of some trees (Fig. 12). Therefore, these terms should be used only at the growth unit annual shoot or axis levels or, at least, the plant level of organization under consideration must be specified when using these terms.

**Hypotony, epitony and amphitony (Fig. 13).** In order to describe the privileged arrangement of lateral axes on a horizontal, curved or slanted parent axis, Troll (1937) defined three modalities of so-called ‘laterale Symmetrie’.

According to this terminology the privileged development of branches on the upper, lateral or basal position of a parent axis is referred to, respectively, as epitony, amphitony or hypotony. Epitony is a very common process in many fruit trees (Costes *et al.*, 2006) and in plants belonging to such architectural models as those of Champagnat, Troll or Mangenot (Hallé *et al.*, 1978); it is also often associated with the survival of old branches in the canopy of old trees (Fig. 13C). Hypotony (Fig. 13A) is frequently marked by a privileged development of lateral axes in the curvature zone of a slanted branch whereas amphitony (Fig. 13B) is frequent on rectilinear horizontal or slightly slanted branches. Hypotony and amphitony may be combined in slanted and curved branches and their incidence in the expansion of lateral branch complexes is of utmost importance in the crown architecture of many woody plants. Finally, it is noticeable that amphitony is a frequent feature in rectilinear branches whereas epitony and hypotony are characterized by the predominant development of lateral axes on the convex side of the curved, downwardly or upwardly orientated branches (Caraglio and Barthelémy, 1997; and compare Fig. 13A, B and C). As highly influenced by axis orientation, these phenomena are frequently combined with the previously described topological arrangement along axes (acrotony, basitony, mesotony) and these combinations have considerable relevance to bud fate according to their topological position and space orientation within a plant crown.

**Morphological differentiation of axes**

The general orientation of a leafy axis and the spatial disposition of its leaves are of major importance in the growth strategy of a plant. Within a single plant, some of these axes are essential in plant skeleton edification; some are aereaerials whereas others may grow underground; some are involved in space exploration whereas others are more related to reproductive dissemination activities or in environment exploitation via photosynthesis. This axis polymorphism is frequent in plants and represents a true morphological differentiation related to meristem expression and activity (Hallé and Oldeman, 1970).

**Orthotropy, plagiotropy and mixed axes (Fig. 14).** On most plants and more evidently in trees, two major types of axes may be distinguished according to their erect or horizontal general orientation. Orthotropy (Fig. 14A; Frank, 1868; Koriba, 1958) refers to axes whose general orientation is vertical and whose symmetry is radial, with leaves in a spiral, opposite or verticillate disposition, and associated lateral branches arranged in all spatial directions. Plagiotropic axes (Fig. 14B; Frank, 1868; Koriba, 1958) have a general horizontal to slanted orientation and a bilateral symmetry owing to leaves (distichous phyllotaxis) and branches being generally arranged in one plane.

Therefore, both kinds of axes are defined not just by their orientation but rather by a set or ‘syndrome’ of features (Edelin, 1984). In addition, some axes may have intermediate features and/or secondary orientation of axes may sometimes occur thus complicating their exact characterization. In many plants, a single meristem may give rise to an axis with mixed properties. Sometimes an axis may have an orthotropic proximal portion and a plagiotropic distal end; the superposition of such ‘mixed axes’ is a distinctive feature of the trunk edification in some plants (Mangenot’s architectural model for instance; Hallé and Oldeman, 1970). Other mixed axes may present the reverse configuration, i.e. a proximal plagiotropic portion followed by a distal orthotropic end. In such cases, the modules are formed by successive hypotonically branching, giving rise to horizontal branched systems named plagiotropy by apposition (Fig. 14C; Hallé *et al.*, 1978; apposing growth of Koriba, 1958; apposing growth of Roux, 1968).
or plagiotropy by substitution (Fig. 14D; Hallé et al., 1978; substituting growth of Koriba, 1958) depending on whether modules are of, respectively, indeterminate or determinate growth.

For a given species, all axes may be of the same kind or several axis types may coexist on the same individual. Different axis types may even coexist at the same foliar axil. On the main stem of Coffea trees for instance, the more distal axillary meristem develops in an immediate plagiotropic branch whereas a more proximal reserve (supernumerary) bud (Varossieau, 1940; Moens, 1963) in the same leaf axil may develop as a delayed orthotropic axis. The coexistence of buds with different fates on the same node leads to many questions regarding the physiological control and genetic determinism of plagiotropy (Tomlinson, 1986, 1987) and more generally branching differentiation.

Short vs. long axes (Fig. 15). In the vegetative aerial part of most woody plants, orthotropy is generally associated with plant skeleton construction and the colonization or exploration of the vertical space, whereas plagiotropy is generally more concerned with exploration and exploitation of the horizontal space and reproductive functions (photosynthesis, flowering).

In many cases, axis differentiation is also related to axis size and very often long and short axes, respectively, specialized in environmental exploration and exploitation (Fig. 15A and B) may be identified in a plant species (Champagnat, 1965; Rivals, 1965, 1966; Kozlowski, 1971; Zimmerman and Brown, 1971). Here again the differentiation of axes and bud fate may be highly specialized and very different structures (i.e. flowers, inflorescences, spines, long axis, etc.) may be found in a single leaf axil and in a precise position, as is the case in several species (see Fig. 15C). In all these cases, however, differentiation of an axis may not be an irreversible process, and according to modifications of internal or external conditions or after architectural traumatism, reversion of axis differentiation is very often possible (see Fig. 15D–F), indicating that shoot differentiation and bud fate are controlled by a whole plant network of correlations (Champagnat, 1961;
Position (terminal vs. lateral) of sexuality and reproductive organs

As architectural studies have historically focused mainly on the vegetative structure of the plant body, reproductive structures are considered as a whole and according to the impact they have on plant growth and branching. Because of their incidence on growth and branching, the lateral or terminal position of reproductive structures will be considered, i.e. whether they result from the transformation of a lateral or terminal meristem, respectively. Readers seeking more precise terminology and descriptions of the structure of reproductive organs or inflorescences in angiosperms should refer to more general (Bell, 1991) or dedicated synthetic works dealing with inflorescence typology or shoot typology according to the arrangement of reproductive or floral elements (Parkin, 1914; Pilger, 1921; Troll, 1957; Briggs and Johnson, 1979; Müller-Doblies and Weberling, 1984; Weberling, 1989; Claßen-Bockhoff, 2000, 2005).

THE CONCEPT OF ARCHITECTURAL MODEL

For a tree species the growth pattern which determines the successive architectural phases is called its architectural
The architectural model is an inherent growth strategy that defines both the manner in which the plant elaborates its form and the resulting architecture. It expresses the nature and the sequence of activity of the endogenous morphogenetic processes of the organism, and corresponds to the fundamental growth programme on which the entire architecture is established. The identification of the architectural model of any given plant is based on the observation of the features belonging to the four major groups of simple morphological features presented above: (1) the growth pattern, i.e. determinate vs. indeterminate growth and rhythmic vs. continuous growth; (2) the branching pattern, i.e. terminal vs. lateral branching vs. no branching, monopodial vs. sympodial branching, rhythmic vs. continuous vs. diffuse branching, immediate vs. delayed branching; (3) the morphological differentiation of axes, i.e. orthotropic vs. plagiotropic vs. axes with mixed morphological and/or geometrical features (with plagiotropic and orthotropic portions); and (4) lateral vs. terminal flowering.

Each architectural model is defined by a particular combination of these simple morphological features and named after a well-known botanist (Fig. 16). Although the number of these combinations is theoretically very high, there are apparently only 23 architectural models found in nature. Each of these models applies equally to arborescent or herbaceous plants, from tropical or temperate regions, and which can belong to closely related or distant taxa.

The reader will find detailed information on each architectural model in Hallé and Oldeman (1970) and Hallé et al. (1978).

Growth patterns defined by the architectural models are genetically determined. Only under extreme ecological conditions is their expression affected by the environment (Hallé, 1978; Barthélémy, 1986; Barthélémy et al., 1995, 1997b, 1999; Grosfeld et al., 1999; Grosfeld, 2002; Stecconi, 2006). Different models can be represented by plants belonging to closely related species (Edelin, 1977; Temple, 1977; Edelin and Hallé, 1985; Vester, 1999; Hallé, 2004). Architectural analysis also shows that some plants frequently exhibit morphological features that are apparently related to two or three models (Hallé and Ng, 1981; Edelin, 1977, 1984; Veillon, 1978, 1980; Grosfeld et al., 1999). These intermediate forms indicate that there is no real disjunction between the models. On the contrary, it must be considered that all architectures are theoretically possible and that there could be a gradual transition from one to another. Among this ‘architectural continuum’ (Edelin, 1977, 1981; Hallé et al., 1978), the models themselves represent the forms that are the most stable and the most frequent, i.e. the most probable biologically. As an architectural model is defined by few and simple morphological features, it gives only an idea of the elementary developmental pattern of a species. Nonetheless, this level of representation of the plant could be well adapted to describe the evolutionary pattern or phylogenetic or taxonomic distribution of plant architecture (Johnson, 2003; Hallé, 2004) and may help to unravel the ecological importance of these patterns (Foresta, 1983; Vester, 1997, 1999).

THE ARCHITECTURAL UNIT

The architectural model represents the basic growth strategy of a plant, and this concept has allowed the definition of a typology of main plant growth strategies. Nevertheless, the characters used in its identification are far too general to describe the complete and precise architecture of a plant. For any given plant, the specific expression of its model has been called its ‘architectural unit’ (Barthélémy et al., 1989, 1991; originally called ‘diagramme architectural’ in French: Edelin, 1977).

The architecture of a plant can be seen as a hierarchical branched system in which the axes can be grouped into categories according to their morphological, anatomical or functional distinctive features (Fig. 17).
The structure and function of each category of axis may be described by a non-limitative series of features. For each of them, the observation of all the architectural characteristics previously described is necessary, but the observations have to be as exhaustive as possible and may concern any elementary level of organization (i.e. metamer, growth unit, annual shoot, module) and any kind of morphological feature (e.g. precise growth direction, phyllotaxis, pre- or neoformation, immediate or delayed branching, form and size of foliar organs, presence, absence and position of sexuality).

The results may be summed up in a table and diagrams that describe and define the specific elementary architecture of each plant, i.e. its architectural unit. Within the context of a general organization, the differences between architectural units are thus represented by the number of categories of axes, their functional and morphological features, and their relative positions.

For each species it has been shown that the number of categories of axes is finite (Edelin, 1977; Caraglio and Edelin, 1990; Caraglio, 1997; Grosfeld et al., 1999; Sabatier, 1999; Grosfeld, 2002; Stecconi, 2006) and generally small (no more than five or six in some Cupressaceae according to the above cited studies). This indicates that the architecture of a fully established branched system, whatever its complexity, can be summarized in terms of a very simple sequence of axes which represents its fundamental organization. In this sequence, leading from axis 1 to the ultimate axis category, following the specific branching pattern, each branch is the expression of a
particular state of meristematic activity and the branch series as a whole can be considered to be tracking the overall activity. In this sense, the architectural unit represents the fundamental architectural and functional elementary unit of any given species.

For structurally complex woody plants in particular, strong axis specialization is frequent: for example, some axes have a more structural and/or exploration function (like the main axis and major branches), and others (short shoots, twigs, etc.) are more concerned with photosynthesis and reproduction. Depending on the species, the differentiation of axes may be strong or not. In some species, axes of different categories have similar features (e.g. ability to flower on all axis categories) whereas some other species have more specialized categories of axes, each presenting very distinctive and exclusive features. Similarly, the relative arrangement of the categories of axes may depend on the degree of differentiation among them, which, in turn, may be related to particular morphological processes such as acrotony. Thus, the categories of axes may or may not be superposed to the notion of branching order (Fig. 18).

According to a species’ developmental pattern, the expression of the architectural unit may be different. Whereas the relative arrangement of categories of axes is generally clear in most woody plants and especially in trees, the total expression of the architectural unit in some herbs or sympodial plants may include the whole succession of modules (Fig. 19).

In a particular species the architectural unit is a very stable and endogenous feature, as shown for several species in which different environmental conditions affect the external form of individuals but change only

**Fig. 18.** Category of axis vs. branching order. The relative arrangement of categories of axes (T, trunk; B, branch; S, short shoot) may (A, i.e. *Araucaria araucana*) or may not (B and C) be superposed to the notion of branching order, in either monopodial (B, i.e. *Acer* sp.) or sympodial (C, i.e. *Platanus* sp.; Caraglio and Edelin, 1990) branching pattern. x, apical mortality; BO

**Fig. 19.** Each category of axis (trunk, T; branch, B) results from the succession of shoots in a monopodial system (A, *Acer* sp., redrawn from Troll, 1937) or the succession of modules in sympodial trees (B, *Ulmus* sp., redrawn from Troll, 1937; dotted lines represent self shed branches; x, apical mortality) or herbs (C, *Encyclia vespa*, from Barthélémy, 1988; dotted sympodial units represent those that are naturally shed for the developmental stage diagrammatically represented).
quantitatively the expression of the species’ endogenous morphogenetic sequence (Fig. 20; Grosfeld et al., 1999).

Finally, for species such as Cupressus sempervirens, for which several ‘forms’ are traditionally well known by foresters and horticulturists (i.e. fastigiated, horizontal and intermediate), it was also shown that there were only minor qualitative differences between the elementary architecture or architectural sequence of these three forms. In this case, the observed phenotypic variability could be explained by the different expressions of several basic morphological and geometrical features such as (1) the relative main stem/branch length, (2) the straightness or gradual straightening-up status of the branches, (3) the insertion angle, (4) homogeneity or heterogeneity of branch types within a single tree and (5) the occurrence, early manifestation and importance of the reiteration process (Fig. 21; Barthélémy et al., 1999).

THE CONCEPT OF REITERATION

Although some plants conform to their architectural unit during their whole life span (Fig. 22), most plants repeat their architectural unit during their development, late in ontogeny, or under particular conditions. Oldeman, (1974) named this process ‘reiteration’ and defined it as a morphogenetic process through which the organism duplicates its own elementary architecture, i.e. its architectural unit. The result of this process is called a ‘reiterated complex’ (Hallé et al., 1978; Barthélémy et al., 1989, 1991) or a ‘reiterate’ (Millet et al., 1998a). Reiteration encompasses several aspects (sprouts, root-suckers, etc.) that have been known incidentally by botanists for a long time. The fundamental interest of this concept resides on it regrouping all these phenomena into a coherent whole, to bring out a common morphogenetic event. Oldeman, (1974) and others (Hallé et al., 1978; Edelin, 1984; Nicolini, 1997; Vester, 1997) hypothesized about the factors triggering this process, and distinguished several types of reiteration.

As first stated by Oldeman, (1974) the reiteration process may involve the expression of the total architectural unit from axis 1 to the most differentiated axis category (‘complete’ or ‘total’ reiteration), or the expression of part of the developmental sequence duplicating only part of the species’ architectural unit (‘partial reiteration’): see Fig. 23A.

Reiterated complexes may originate from dormant meristems and reiteration in this case is called ‘proleptic’ or ‘delayed’. By contrast, reiteration may result from a shift in the functioning of the apical meristem of a growing shoot that will finally produce a ‘less differentiated structure’, i.e. a branch apex that after some time of functioning gives rise to a ‘supernumerary trunk’. In this case, the reiteration is described as ‘sylleptic’ (or better ‘immediate’) or ‘reiteration by dedifferentiation’ (Fig. 24). Either of these two types of reiterations may be qualified as total or partial.

Reiteration at first was considered as an opportunistic (non-automatic) process (Oldeman, 1974; Hallé et al., 1978). ‘Opportunistic reiteration’ may today be considered as any kind of reiteration linked to the individual history of each tree and may have two main origins: (1) ‘adaptive reiteration’ is a response to an increase in resource levels whereas (2) ‘traumatic reiteration’ is a response of a plant after it has been damaged and lost a major part of its structure (Fig. 25). More recent investigations (de Castro e Santos, 1980; Edelin, 1984; Sanoja, 1992; Nicolini, 1997; Sabatier, 1999; Vester, 2001; Grosfeld, 2002; Stecconi, 2006), however, have

**Fig. 20.** Variation in crown physiognomy and architecture in relation to environmental conditions at the time of architectural unit expression in Araucaria araucana. (A) In forest stands, the mature tree expressing its architectural unit has a 15–20-m-high trunk which bears at its top a large conical crown composed of up to 20 tiers of living branches. (B) In full sun and with favourable soil and precipitation conditions, the tree has, at first cone production, a typical pyramidal crown. The trunk is 6–8 m high and most branches are alive. (C) In full sun and poor soil and precipitation conditions, the first production of cones may occur in a tree no more than 4 m high. The crown has a typical ‘umbrella form’ and most of the branches are alive (Grosfeld et al., 1999). Black arrows indicate terminal female cones.
demonstrated that, beside these cases of opportunistic reiteration, the same process of repetition may be involved in the inherent growth pattern of a species and occur automatically during plant development after a definite threshold of differentiation (Fig. 26). This latter case is a common feature of tree development and crown construction and is referred to as ‘automatic’ (Edelin, 1984) or ‘sequential’ reiteration (Nicolini, 1997).

As already noted, the development of a plant conforming to its model implies the notion of a differentiation sequence in the activity of the whole set of meristems of the plant (see also following sections). In the case of opportunistic reiteration (cf. Fig. 23) the occurrence of reiterated complexes seems to be a move backwards within the plant’s developmental sequence, i.e. a real ‘dedifferentiation’. A supernumerary trunk (or branch), resulting from the transformation of a growing branch (Fig. 24, ‘immediate reiteration’), or from the development of a branch from a dormant meristem (Fig. 23, ‘delayed reiteration’), implies that the plant expresses again the juvenile growth pattern of the organism developed from seed. This is well illustrated in cases of regeneration in which, when a trunk is cut, sprouts resembling young trunks are formed from the stump, whereas reiterated complexes that develop after a branch has been damaged have an architecture similar to that of this branch. More generally speaking, it has been demonstrated that, in this case, the real degree of dedifferentiation of a reiterated complex depends on the location of the reiterated complex in the whole plant architecture and on the ontogenetic stage of the plant at the moment reiteration occurs (Barthélemy, 1988; Fig. 23B, C).

‘Automatic’ or ‘sequential’ reiteration has been intensively studied in the last two decades (Edelin, 1984; Caraglio and Edelin, 1990; Barthélemy et al., 1991;...
plexes (in black) according to their location on the tree in (B) _Symphonia globulifera_ (from Barthe´le´my, 1988) and in (C)

At the top of the tree and in the most peripheral part of the crown, pau-

At the top of the tree and in the most peripheral part of the crown, paup-

Whatever their qualitative or quantitative nature these vari-

Whatever their qualitative or quantitative nature these vari-

At the top of the tree and in the most peripheral part of the crown, paup-

Fig. 23. Opportunistic reiteration and structure of the reiterated com-

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**LEVELS OF ORGANIZATION, REPETITION PHENOMENA AND SEQUENCES OF DIFFERENTIATION**

Since the pioneering work of von Goethe (1790), observa-

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Durand, 1997; Nicolini, 1997; Vester, 1997; Sabatier, 1999; Grosfeld, 2002; Stecconi, 2006) and proved to be a

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environmental factors may ‘modulate’ these sequences of differentiation (Fig. 20), it was shown (Barthélemy et al., 1995, 1997a, b; Nicolini 1997; Sabatier, 1999; Claßen-Bockhoff, 2000; Heuret et al., 2000; Guérard et al., 2001) that they almost never (except probably in extreme conditions) modify the inherent morphogenetic and ontogenetic constructional rules of plant organization.

Fig. 24. (A) Total immediate reiteration (S.R.) expressed on the distal part of a branch of Araucaria araucana. (B) Successive immediate and partial reiterated complexes on a branch of Austrocedrus chilensis (from Grosfeld, 2002). A1, main stem; A’2, first order of reiteration of A2 category of axis; A”2, second order of reiteration.

Fig. 25. (A) Damaged tree crown of an unidentified tropical tree comprising a part of the initial crown (C) and a set of reiterated complexes (C.R.) on the broken part (x). (B) Adaptive reiterated complexes (C.R.) can occur following local structural changes in the crown of an individual of Fraxinus excelsior (Barthélemy et al., 1997a) as caused by traumatism along the main stem (x); compare with Fig. 26.
THE NOTION OF ‘MORPHOGENETIC GRADIENTS’

In the last two decades, coupled with precise morphological observations, architectural analyses of several plant species (Caraglio and Edelin, 1990; Barthélémy et al., 1992, 1995; Drénou, 1994; Nicolini and Caraglio, 1995; Caraglio and Barthélémy, 1997; Nicolini, 1997, 1998, 2000; Gleißner, 1998; Puntieri et al., 1998, 1999, 2000, 2001, 2002a, b; Genoyer et al., 1999; Nicolini and Chanson, 1999; Sabatier and Barthélémy, 1999, 2001a, b; Sabatier et al., 1999, 2003a, b; Claßen-Bockhoff, 2000; Heuret et al., 2000; Nicolini et al., 2000; Souza et al., 2000; Stecconi et al., 2000; Guérard et al., 2001; Passo et al., 2002) revealed that, under given environmental conditions, the structure and features of a particular elementary botanical entity (metamer, growth unit, annual shoot) are predictable and strongly dependent on both (1) its topological location in the comprehensive architecture of a plant and (2) the ontogenetic stage of the organism.

At the level of the whole plant, the ‘morphogenetic gradients’ notion was defined (Barthélémy et al., 1997a) in order to take into account the intrinsic organization rules of plant structure and was shown to be a powerful concept (Prusinkiewicz et al., 2001) to explain the observed structure and series of modifications of botanical entities during the development as a whole as they are associated with either the first successive steps of the ontogeny or the ageing of axes or plants. These gradients were respectively called ‘base effect’, or ‘drift’ (Barthélémy et al., 1997a) and take into account the respective ontogenetic increase or decrease in several parameters along the main stem of any plant, or even along lateral axes of branched plants. They encompass several phenomena that have been known and described for a long time but that very often were considered or defined independently or separately according to the observer’s interest (plant structural edification for botanists, main stem height and radial growth for foresters) and the phylogenetic status (seed plant or not, actual or fossil plant, monocotyledons or dicotyledons, etc.) or the particular growth pattern of the plant, or plant group (i.e. monopodial or sympodial, herbaceous annual or woody perennial plants) under study.

According to the specific complexity and/or growth pattern, these gradients may be described using different parameters. Among these parameters are the size, nature, and/or internal and external structure of the successive metamers, growth units, annual shoots or modules in sympodial plants or even appendages and sibling lateral axes if any (Fig. 31), but also on the growth rate and the various portions of the sigmoidal growth curve of all continuously growing plant in which morphogenetic gradients will only concern the successive metamers (and their individual constitutive parts) whereas it will be at a maximum in a polycyclic, branched and reiterated, rhythmically and sympodially growing plant where these gradients will reflect, at several levels, the complex topological and geometrical structural nested organization.

Some of these gradients are very widespread in the plant kingdom and have a very comprehensive and general expression during the development as a whole as they are associated with either the first successive steps of the ontogeny or the ageing of axes or plants. These gradients were respectively called ‘base effect’, or ‘drift’ (Barthélémy et al., 1997a) and take into account the respective ontogenetic increase or decrease in several parameters along the main stem of any plant, or even along lateral axes of branched plants. They encompass several phenomena that have been known and described for a long time but that very often were considered or defined independently or separately according to the observer’s interest (plant structural edification for botanists, main stem height and radial growth for foresters) and the phylogenetic status (seed plant or not, actual or fossil plant, monocotyledons or dicotyledons, etc.) or the particular growth pattern of the plant, or plant group (i.e. monopodial or sympodial, herbaceous annual or woody perennial plants) under study.

According to the specific complexity and/or growth pattern, these gradients may be described using different parameters. Among these parameters are the size, nature, and/or internal and external structure of the successive metamers, growth units, annual shoots or modules in sympodial plants or even appendages and sibling lateral axes if any (Fig. 31), but also on the growth rate and the various portions of the sigmoidal growth curve of all
plants and axes (Pressler, 1865; Pardé and Bouchon, 1988) or even on the ‘rhythms of primary thickening growth’ (Kaplan, 2001) described by various authors (‘Erstarkungswachstum’ of Troll and Rauh, 1950; ‘establishment growth’ of Tomlinson and Zimmerman, 1966; Tomlinson and Esler, 1973; ‘epidogenesis’ and ‘apoxogenesis’ of Eggert, 1961, 1962; Daviero et al., 1996; Soria and Meyer-Berthaud, 2004). These rhythms are often associated with the obconic shape of primary structures (Martinez and De La Sota, 2001; Hueber and Galtier, 2002; Isnard et al., 2005) during the first stages of shoot (or even plant) ontogeny and/or with the diameter of the shoot apex (Steeves and Sussex, 1989). Whatever their specific and particular expression and because of their very wide occurrence in any actual or fossil plant, these gradients or ‘rhythms of primary thickening growth’ may be considered as one of the few fundamental principles of plant shoot organization (Kaplan, 2001).

‘Base effect’ and ‘drift’ are commonly expressed at the axis level and thus affect the main axis of any plant and all the lateral axes of branched ones. According to these gradients, the exact structure of a lateral axis depends on its topological and ontogenetic position on the parent axis (Fig. 30). The effect of ‘branching order’ may be superimposed on these previous gradients at a whole plant level. In general terms, the higher the branching order of an axis, the higher its degree of differentiation. At a more local scale in the architecture of a branched plant or system and according to the expression of the specific branching pattern, gradients, linked to shoot structure and precise branching pattern (acrotony, basitony, mesotony) or linked to shoot spatial orientation and/or geometry (hypotony, epitony, amphitony), may be superimposed on previous ones, which explains the observed heterogeneity of axes or shoots in close topological positions.
Because of its duplicative and repetitive nature, the reiteration process reproduces the morphogenetic gradients of the non-reiterated parent plant (Fig. 30) and the structure of the reiterated complexes resembles that of the whole plant (Fig. 23B, C; Barthélémy, 1988).

As previously stated, all these morphogenetic gradients have a strong intrinsic basis and allow, in a given environmental condition, the prediction of the precise structure of a particular elementary entity according to its precise architectural position and to the developmental stage of the plant. Their expression, however, may be modulated by environmental or technical factors. Where constant but different conditions prevail during the whole ontogeny, this may lead to very different final structures as development in each condition may be related to a particular ‘reading’ of the intrinsic sequence of differentiation or in other words a particular ontogenic trajectory (Fig. 17). In fluctuating conditions, in case of a trauma or according to localized and temporally limited perturbations resulting from environmental or technical causes, knowledge of these gradients and the possible a posteriori identification of their local or partial alteration or modification and the induced local perturbation of the sequence of differentiation may also serve as a basis for a very precise diagnosis of a plant’s particular history (Fig. 25B).

THE NOTION OF ‘PHYSIOLOGICAL AGE OF A MERISTEM’

The notion of morphogenetic gradients illustrates and takes into account the commonly observed situations where, for a given plant species or even individual, (1) at a given time and/or for a given stage of development, homologous botanical entities with very different features (or ‘states of differentiation’) coexist on the same individual (e.g. short vs. long shoots or reproductive vs. vegetative shoots; for instance see Figs 19, 21, 23 and 27) whereas, by contrast, (2) similar elementary botanical entities with the same morphological features (e.g. short floriferous shoots for instance; e.g. see Figs 15 and 29) may be encountered for very different plant ages or stages of development.

These observations complemented by others led to the definition of ‘physiological age of a meristem’ (Barthélémy et al., 1997a) that may generally be characterized and defined by a particular combination of several morphological, anatomical and/or functional attributes of a given botanical entity derived from this meristem (Fig. 32).

This physiological age of a meristem is thus determined, a posteriori, by the morphological analysis of the elementary botanical entity it produces, and may not be an intrinsic property of this meristem itself but the result of all morphogenetic interactions between plant parts that are related to differentiation processes and result in the expression of morphogenetic gradients.

As previously discussed, the physiological age of a meristem (or elementary botanical entity) depends on its precise location in the plant architecture and on the stage of development of the organism, and its expression may be modulated by environmental factors. The functioning of a meristem, or the elementary botanical entities it produces, can thus be characterized by three different ages (Barthélémy et al., 1997a; Fig. 33):

1. the calendar or chronological age corresponds to the period (i.e. year, month, week or day of formation for instance) in which the elementary botanical entity has been edified (metamer, growth unit or annual shoot according to specific and relevant growth pattern and plant complexity);
2. the ontogenetical age refers to the elapsed time after seed germination (the ontogenetic time unit considered may be a year, a day or a growth cycle according to...
(3) The physiological age of a meristem relates to the degree of differentiation of the structures it produced. It may be estimated \textit{a posteriori} by a non-limitative series of qualitative and quantitative criteria. For example, the short axes of many trees are typical features of ‘physiologically aged’ structures: growth units are short, bear flowers and have a short lifetime. These highly differentiated axes may be considered as ‘physiologically old’ whatever their moment of appearance. By contrast, main axes consisting of vigorous growth units and/or annual shoots may be considered as ‘physiologically young’ products and generally appear only in the young tree.

The identification for a botanical entity of these three ages is fundamental in order to understand the comprehensive architecture of a plant or even its plasticity, i.e. the effects of the environment on its development and structure. It permits the precise characterization of all elementary levels of organization within the more integrative individual architecture and allows a precise multilevel description of plant architecture and organization.

\textbf{CONCLUSIONS}

Any plant may be recognized by its general form, but this physiognomy is always the result of a very precise structure that underlies the existence of a strong inherent organizational pattern in plant architecture construction.

Architectural studies that have been carried out for some 30 years have led to the definition of several concepts and notions that provide powerful tools for studying plant form and intrinsic morphogenetic and ontogenetic
and architectural plasticity (Grosfeld et al., 1999; Nicolini et al., 2000; Guérard et al., 2001; Stecconi, 2006), technical practices (Caraglio et al., 2000; Sabatier et al., 2000; Leroy and Caraglio, 2003), architectural and genetic variation, plant breeding and selection (Barthélemy et al., 1999; Corradi et al., 2002; Sabatier et al., 2003a) and landscape management (Auclair et al., 2001), among other issues.

Based on this botanical background and with the development of high-power computers, a concomitant and complementary computational and numerical approach to the modelling and computer visualization of plant architecture has gained importance in recent decades. Mathematical models of growth, branching and architecture have been developed on the basis of qualitative botanical knowledge, e.g. models and quantitative data on the demography, size and positions of plant components (Godin et al., 1997, 1999; Godin and Caraglio, 1998; Ferraro and Godin, 2000, 2003; Godin, 2000; Guédon et al., 2001; Costes and Guédon, 2002; Heuret et al., 2002, 2003; Costes et al., 2003; Durand et al., 2005; Ferraro et al., 2005; Godin et al., 2005). This new disciplinary approach offers new possibilities to understand or quantify the genetic basis or regulation of plant architecture (Barbier de Reuille et al., 2005, 2006; McSteen and Leyser, 2005; Mündermann et al., 2005; Wang and Li, 2005) and will probably renew various aspects of developmental biology studies (Turnbull, 2005).

Results of these mathematical models are used to simulate plant development and architecture (de Reffye and Houllier, 1997). In these simulations much emphasis has been given to the changes in the physiological age of plant components and in the three-dimensional structure during plant development (Bouchon et al., 1997; Yan et al., 2003; Zhao et al., 2003). This methodology may be used for any kind of plant (Barczi et al., 1997) and has also been successfully used to simulate root systems (Jourdan and Rey, 1997a, b, c; Danjon et al., 1999a, b, 2005; Dupuy et al., 2005a, b, c, 2006). It can also be used to evaluate developmental changes such as growth responses to controlled environments, aerial biomass, volume prospected in soil by roots, radiant energy transfer, retrodiffusion of the canopy for calibrating remote-sensing techniques, and the mechanical status of a tree (de Reffye and Houllier, 1997; Castel et al., 2002; Hu and Jaeger, 2003; Soler et al., 2003; Picard et al., 2004; Chenu et al., 2005; Sellier et al., 2006). Finally, the models may be linked to eco-physiological models, leading to structural functional models (Godin and Sinouquet, 2005) or process-based architectural models, i.e. morphogenetic models of tree architecture directly driven by eco-physiological processes, with possible applications in environmental and genetic control of morphogenesis in crops and phenotypic plasticity modelling and simulation (de Reffye et al., 1997; Yan et al., 2004; Dingkuhn et al., 2005; Guo et al., 2006).

Because of its economic importance for agronomic production, plant reproduction or vegetative propagation, acquisition of reproductive capacities during ontogeny has been extensively studied (Goebel, 1900; Passecker, 1944, 1958; Chouard, 1950; Rémy, 1951; Stokes and Verkerk,
Fig. 31. (A) Shape and size of successive leaves along a stem of a herbaceous *Senecio* sp. (redrawn from Baillaud and Courtot, 1955). (B) Organization of the two last successive annual shoots of the main stem of *Cedrus atlantica* during its life and (C) the corresponding global tree structure (from Sabatier and Barthéléméy, 1995).

Fig. 32. Diagrammatic representation of trends in the value of some morphological parameters according to annual shoot rank of successive annual shoots of branches borne on the same ontogenetical age parent shoots of the main stem of a set of 22-year-old dominant to co-dominant *Pinus pinaster* individuals grown on a same stand in the south-west of France (unpublished data; Coudurier et al., 1995; Heuret et al., 2006), where *a* is the mean annual shoot length (in percentage of maximum length), *b* is the percentage of polycyclic shoots, *c* is the percentage of branched shoots, and *d* is the percentage of shoots with male cones. For each shoot rank the particular combination of the value of the morphological parameters allows a strict characterization of the physiological age of meristem activity that produced it.

Fig. 33. Diagrammatic representation of a theoretical plant whose elementary botanical entities produced by the meristems (for instance annual shoots, represented by rectangles) may encompass four different physiological ages (a–d). The ‘plant’ grows from seed in four steps (possible ontogenetic ages 1–4), each step corresponding to one year of growth (n – 3 to n). It is hypothetically represented as growing in ‘good’ environmental conditions (where the main stem expresses the four possible physiological ages successively and in relation to successive four ontogenetic ages and where branches fully express morphogenetic gradients as shown in Fig. 30: case ‘A’), or in suppressed condition (as for the growth of *Araucaria araucana* in Fig. 20; case ‘B’). Each entity is characterized by a trinom – where the first element represents ontogenetic age (1–4), the second element represents calendar age (n – 3 to n) and the third element represents physiological age (a–d) – that allows us to understand the structure of each entity in the whole architecture of the plant (represented in A and B 4 years after germination).
The general description of plant parts during the whole ontogeny. In the latter cases, the idea of heterogeneity of plant parts either at a given stage or level is not sufficient to take into account the complete levels of organization we have demonstrated that metamer is known that it may be strongly modified by environmental factors (Koch, 1873; Carrière, 1880; Hochtetter, 1880; Beissner, 1888; Cockayne, 1912; Frost, 1938; Webber and Batchelor, 1948; Ashby, 1949; Doorenbos, 1954; Schaffalitzky de Muckadell, 1954, 1959; Brink, 1962; Trippi, 1963; Wardle, 1963; Greenwood and Atkinson, 1977; Poethig, 1990, 2003; Jones, 1999; Kaplan, 2001) and these changes may be simultaneous with the appearance of reproductive capacity during development. At the whole individual level, plant development may thus be viewed as a succession of phases, each characterized by a set of several features or parameters. Under given conditions, the duration of these phases may be more or less fixed for a species (Wareing, 1959; Zimmerman, 1972; Hackett, 1985) and it is thus possible to identify a mean specific age in which a plant will be able or unable to express a particular feature. According to plant species, this ‘chronological’ (Ritterbusch, 1990) or ‘calendar’ (Gatsuk et al., 1980) age may be expressed in days or years after germination and it is known that it may be strongly modified by environmental factors (Doorenbos, 1954; Zimmerman, 1972; Gatsuk et al., 1980). At a given moment, a plant may thus be characterized not only by its ‘chronological’ or ‘calendar’ age but also by a set of biological criteria that indicates its ‘stage’ of development variously referred to as ‘biological age’ (Levin, 1966; Roloff, 1989; Gleißner, 1998), ‘physiological age’ (Robbins, 1957; Schaffalitzky de Muckadell, 1959; Grubb, 1977), ‘ontogenic age’ (Passecker, 1977) or ‘age state’ (Uranov, 1975; Gatsuk et al., 1980). The ‘physiological age’ of the individual may merge with the metamer level and may be defined or even quantified by the features or parameters of the successive constitutive metamers or leaves in small, herbaceous monocalous continuously growing plants, as variably demonstrated (Ashby, 1949; Baillaud and Courtot, 1955; Poethig, 2003; Fig. 31A), because the whole plant structure corresponds to an elementary botanical entity. In more complex plants with several nested levels of organization we have demonstrated that metamer level is not sufficient to take into account the complete heterogeneity of plant parts either at a given stage or during the whole ontogeny. In the latter cases, the idea of a ‘physiological age’ better corresponds to the growth unit and/or sympodial unit and/or annual shoot levels (Fig. 31B). The general description of plant parts heterogeneity thus implies the use of this notion at the meristem level and the pertinent elementary levels it produces in order to take into account plant architecture diversity. Coupled with the description of ‘morphogenetic gradients’, the notion of physiological age of a meristem (as applied to the pertinent level of organization) may allow a generic description of the multiple and diverse expressions of ‘ontogenetic contingency’ (Diggle, 1994, 1997) observed in the plant kingdom (Kaplan, 2001). Furthermore, as these notions allow a multilevel description of the influence of the environment on several features and their expression along ontogeny and according to architectural position, they may provide a powerful tool to separate ontogenetic variations from environmental plasticity or to separate traits or ontogenetic trajectories plasticity (Wu and Stettler, 1998; Huber et al., 1999; Wu and Hinckley, 2001; Alpert and Simms, 2002; Diggle, 2002; Novoplansky, 2002; Wright et al., 2002; Sachs, 2004; de Kroon et al., 2005; Wolfe and Mazer, 2005).

Despite the frequently stressed importance of considering plant morphology and/or architecture to understand their phylogeny, very few studies actually describe the link between phylogeny and architecture (Sanoja, 1992; Johnson, 2003; Hallé, 2004) and here also we probably require more data and a multilevel consideration of plant structure.

Finally, the dynamic, multilevel and comprehensive approach of plant form and ontogeny have proved to be fruitful but the number of questions raised shows that some of the answers will also, we hope, come from a multidisciplinary approach and by the combination of different approaches to the same plant object.

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