Summary

Vacuoles are present in the cells of plants, fungi, protists and animals. However, their importance is most pronounced in plants due to the multiplicity of functions and the dimensions of vacuoles in plant cells. An interesting, and relatively recently discovered aspect of plant vacuoles is that some cell types have multiple vacuoles with very different content and functions. This raises questions how multiple vacuoles can be formed in a single cell and how proteins are directed to the one or the other vacuolar compartment.

The identification of vacuolinos, a novel type of additional vacuole that exists besides the anthocyanin containing central vacuole in petal epidermal cells, provided an excellent model system to study the formation and functional differentiation of vacuoles, but not without raising new issues and questions. Each chapter of this thesis is an immersion in the biology of vacuoles, with special emphasis on the biogenesis, maturation and physiology of vacuolinos. Vacuoles are here investigated by a combination of molecular genetics, cell biology and bioinformatics. The results that were obtained provide new insights into the mechanism(s) of regulation of vacuolar pH in extremely acidified cells, the biogenesis and biological function of vacuolinos and the mechanism(s) by which proteins transport from vacuolinos to the central vacuole is controlled.

In Chapter 1, protein sorting pathways and the biogenesis vacuole in plants and fungi and lysosome in animals, as well as the formation of functionally distinct vacuoles in a single plant cells are reviewed. Comparative analysis of the genes that were shown to be involved in the biogenesis of these compartments suggests both conservation and divergence of the components of the machinery that drives
biogenesis of and protein trafficking to these compartments during evolution in plants, fungal and animal lineages.

The analysis of petunia mutants impaired in vacuolar acidification in the petal epidermis, revealed that vacuolino formation is controlled by the same regulatory network that governs hyper-acidification of the central vacuole where anthocyanins are stored. The transcription factors PH4 (encoding a MYB protein), AN1 (a bHLH), PH3 (a WRKY) and AN11 (a WD repeat protein) together form the MBWW transcription complex. This complex activates the expression of \( PH1 \) and \( PH5 \), encoding two vacuolar P-ATPases, which hyperacidify vacuoles in petal epidermal cells, conferring a reddish color to the petals, in contrast to the bluish color of mutants affecting this pathway. This very same complex regulates a set of downstream target genes involved in vacuolino biogenesis.

In **Chapter 2**, the machinery responsible for the hyper-acidification of vacuoles in the flesh of *Citrus* fruits is characterized, and shown to be the very same one described for petunia petals. \( CitPH1 \) and \( CitPH5 \), are expressed in sour lemons, oranges, pummelos and rangpur lime fruits, while their expression is strongly down-regulated in sweet-tasting varieties, which can be traced back to mutations affecting the MBWW complex. Because of the proven role of the MBWW complex in both vacuolar acidification and vacuolino biogenesis, the identification of all components of MBWW complex and the demonstration of its involvement in the expression of \( PH1 \) and \( PH5 \) in *Citrus*, strongly suggests the existence of vacuolino in fruits cells. These considerations open the way to investigate the distribution of vacuolinos in different tissues and different species.

Since their identification, vacuolinos, were shown to be intermediate stations for a range of different proteins on their way to the central vacuole, but the biological
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function(s) of these organelles remained largely mysterious, as mutant flowers that lack vacuolinos (e.g. \textit{an1}, \textit{ph4}) do not display any obvious macroscopically visible defects. In specific genetic backgrounds, containing a dominant allele of the \textit{FADING (FA)} locus, loss-of-function \textit{ph4} mutations cause besides the lack of vacuolinos in petal epidermal cell, also the degradation of anthocyanin pigments in the central vacuole and “fading” of the petal color after bud opening. Although the genetics of this phenomenon was described already in the 1980s, the underlying mechanism(s) remained obscure. \textbf{Chapter 3} describes the isolation and characterization of the \textit{FADING (FA)} gene, the key factor in the fading process. \textit{FA} encodes a member of the ferric reductase oxidase (FRO) group of the family of metalloreductase enzymes, and is expressed in petals as well as various vegetative tissues, like leaves and stems.

In petal epidermal cells of mutants for the MBWW complex (\textit{an1}, \textit{ph3} and \textit{ph4} mutants), \textit{FA} localizes on the tonoplast of the central vacuole where the anthocyanin molecules are stored. These cells lack vacuolinos and \textit{FA} reaches the central vacuole where it (probably indirectly) destabilizes anthocyanins. In In petal epidermal cells of wild type Petunia lines, which contain vacuolinos, GFP and RFP-tagged \textit{FA} protein moves to vacuolinos, just like other vacuolar proteins. Within vacuolinos, \textit{FA}-GFP (or at least the GFP tag) is released from the membrane and now accumulates in the vacuolinos lumen, which may explain why \textit{FA}-GFP does not move on from vacuolinos to the central vacuole, as other vacuolar proteins do, and thus cannot trigger the degradation of anthocyanins in there. In the petals epidermis of \textit{ph4} and \textit{ph3} mutants, by contrast, vacuolinos are abolished and \textit{FA}(-GFP) now reaches the vacuoles, like it does in leaf cells, to trigger the degradation of anthocyanins in there. These findings suggest that vacuolino may serve as a gatekeeper/check point that allows certain proteins to reach the central vacuole, while keeping others away from the central vacuole. Moreover, the new insights in the control of flower (and fruit) pigmentation via the regulation of anthocyanin...
degradation, which may be helpful to generate crop varieties with new flower colors or enhanced dietary value of fruits

Previous work showed that the MBWW transcription factor complex is essential for the formation of vacuolinos. MBWW was thought to act indirectly, most like through the activation of downstream genes encoding proteins that are directly involved in membrane trafficking and vacuolino formation. However, the pathway by which vacuolinos are formed and the (MBWW regulated) genes and proteins that are involved remained unknown. Chapter 4 presents the identification of the first target gene, RAB5a, of the MBWW complex that is required for the biogenesis of vacuolinos. RAB5 is a member of the RAB family of small GTPases, many of which have crucial roles in specifying the “identity” of membranes and the docking and recognition vesicle and target membranes. The identification of RAB5a, a homolog of metazoan RAB5, showed that plant RAB5s are evolutionary more diverse than previously thought and consist of three distinct phylogenetic clades, represented in petunia by RAB5a, RAB5a1, and RAB5a2 plus the homologous (and well-studied) Arabidopsis proteins RHA1 and ARA7. The presented data show that the RAB5a gene is activated by the MBWW complex and that the RAB5a protein is needed to facilitate the fusion of a subpopulation of prevacuolar compartments (PVCs) to form vacuolinos. Interestingly, RAB5a1 and RAB5a2, localize to different cellular compartments than RAB5a, and cannot functionally substitute RAB5a. This highlights the pivotal role of the RAB5a GTPase in vacuolino formation and provides insights into the diversification of distinct vacuolar compartments during evolution, through duplication and neofunctionalization of genes encoding key regulators of vesicle trafficking.

The abolishment of vacuolinos in rab5a mutants, and the formation of enlarged vacuolino induced by expression of RAB5a from the constitutive 35S promoter,
affect in Petunia the dimension and shape of petal epidermal cells. Such effect on epidermal cells might alter petal coloration and even pollinator preference, and a deeper analysis of the consequences of RAB5a activity on cell structure, might enrich our understanding of the diverse biological function of vacuolino.

Mutants resulting in petal color shift towards blue, defined seven so-called PH loci in petunia. The loci control hyper-acidification of the central vacuoles in the petal epidermis and thereby the color displayed of the anthocyanins pigments in there, by altering protonation state at different pH. Of these seven PH loci, six had been molecularly analyzed previously and their biological functions described. However, PH7, remained so far uncharacterized. In Chapter 5, comparative analysis of RNA-seq data generated from PH7 wild type, ph7 transposon unstable mutant and ph7 stable mutant have successfully identified the PH7 gene. PH7 encodes a monosaccharide sugar transporter that is highly homologous to the Arabidopsis ERD6-like6 glucose/proton symporter residing on tonoplast. Mutations in different regions of PH7 result in distinct defects in the vacuolino pathway, blocking either the formation of vacuolinos (and possibly downstream steps), or the trafficking of proteins from vacuolinos to the central vacuole. This indicates that PH7 is required at multiple steps along the vacuolino pathway, and that it facilitates the formation of vacuolinos and the subsequent vacuolino-central vacuole trafficking through distinct protein domains and mechanisms. Although, further research is needed to unveil in detail the mechanism underlying the petal color change in ph7 mutants and the role of this transporter in the physiology of vacuolino membranes, the current findings present evidence that supports previous hypotheses/suggestions that membrane transporters may have additional functions in mediating recognition or fusion of vesicles and target membranes during vesicle and proteins trafficking.
Samenvatting

Vacuolen zijn organellen die voorkomen in cellen van planten, schimmels, protisten en dieren. De vacuolen in plantencellen zijn het meest bekend, vanwege de veelheid aan functies en de enorme afmetingen van vacuolen in plantencellen. Een interessant en relatief recent ontdekt aspect van plantenvacuolen is dat sommige celtypen meerdere vacuolen hebben met zeer verschillende inhoud en functies. Dit roept vragen op hoe meerdere vacuolen kunnen worden gevormd in een enkele cel en hoe eiwitten worden gericht naar het ene of het andere vacuolaire compartiment.

Vacuolinos, zijn kleine additionel vacuolen recent beschreven in epidermale cellen van bloembladen waar ze voorkomen naast de grote centrale vacuole die de anthocyaan bloemkleurpigmenten bevat. Deze additionele vacuolen vormen een uitstekend modelsysteem om de biogenese en functionele differentiatie van vacuolen te bestuderen. Tegelijkertijd werpen deze vacuolinos ook nieuw vragen op, onder andere met betrekking tot hun biologische functie. Elk hoofdstuk van dit proefschrift is een reis door de biologie van vacuolen, met speciale nadruk op hun biogenese, rijping en fysiologie. Vacuolen worden hier onderzocht door een combinatie van methodes uit de moleculaire genetica, celbiologie en bio-informatica. De verkregen resultaten verschafften nieuwe inzichten in de mechanismen die ten grondslag liggen aan de regulatie van vacuolaire pH regulatie in extreem verzuurde cellen, de biogenese en biologische functie van vacuolinos en het mechanisme waardoor eiwitten van vacuolinos naar de centrale vacuole worden gestuurd.

In hoofdstuk 1 worden eiwitsorteerroutes en biogenese van vacuolen in planten en schimmels, en lysosomen bij dieren besproken, alsmede de vorming van meerdere functioneel verschillende vacuolen in een enkele plantencel. Analyse en vergelijking van de genen waarvan is aangetoond dat ze betrokken zijn bij de biogenese van deze
compartimenten suggereerde zowel conservering als divergentie tijdens de evolutie van planten, schimmels en dieren van de componenten van de factoren die de biogenese van vacuolen en lysosomen en de sortering van eiwitten naar deze compartimenten reguleert.

De analyse van petunia mutanten met defecten in verzuring van de vacuolen in de epidermis van kroonbladeren (petalen), liet zien dat de biogenese van vacuolino’s wordt gecontroleerd door hetzelfde regulatoire netwerk dat de sterke verzuring regelt van de centrale vacuole. De transcriptiefactoren PH4 (coderend voor een MYB-eiwit), AN1 (een bHLH eiwit), PH3 (een WRKY eiwit) en AN11 (een WD-repetal eiwit) vormen samen het MBWW-transcriptiecomplex. Dit complex activeert de expressie van *PH1* en *PH5*, coderend voor twee vacuolaire P-ATPases die vacuolen in epidermale cellen van bloemblaadjes verzuren, en een roodachtige kleur veroorzaken in de bloemblaadjes, in tegenstelling tot de blauwachtige kleur van mutanten die deze route beïnvloedden. Ditzelfde complex reguleert ook een aantal genen die betrokken zijn bij vacuolino-biogenese.

**In hoofdstuk 2** wordt de machinerie gekarakteriseerd die verantwoordelijk is voor de extreme verzuring van vacuolen in *Citrus* vruchten en wordt aangetoond dat deze machinerie dezelfde te zijn als die beschreven voor petunia bloemblaadjes. *CitPH1* en *CitPH5* komen sterk tot expressie in sterk zure citroenen, sinaasappels, pummelo’s en rangpur limoenen, terwijl hun expressie sterk is verlaagd in niet-zure vruchten met een zoete smaak. Dit bleek het gevolg van mutaties die de expressie of de activiteit het MBWW-complex beïnvloeden.

In petunia bloemen is het MBWW-complex essentieel voor de expressie van genen betrokken bij vacuolaire verzuring en voor genen die een rol spelen in de vorming van vacuolino’s. Analyse van publieke transcriptoom data wees uit dat in zoete sinaasappels, niet alleen expressie van *CitPH1* en *CitPH5* is verlaagd, maar ook de
expressie van *Citrus* homologen van genen die in petunia essentieel zijn voor de vorming van vacuolinos, zoals *RAB5a*. Dit suggereert vacuolinos ook voorkomen in ander weefsels dan de bloembladepidermis. Deze overwegingen openen de weg om de aanwezigheid en functie(s) van vacuolino in verschillende weefsels en verschillende soorten te onderzoeken.

Vacuolinos zijn kleine vacuole-achtige compartimenten die voorkomen naast de grote centrale vacuole in epidermale cellen van bloemblaadjes. Sinds hun identificatie, bleken vacuolinos tussenstations te zijn voor een reeks van verschillende eiwitten op weg naar de centrale vacuole, maar de biologische functie(s) van deze organellen bleef grotendeels mysterieus. Eén reden is dat bloemen die vacuolinos missen als gevolg van een mutatie (bijv. *an1*, *ph4*) geen duidelijke macroscopisch zichtbare defecten vertonen. Echter in specifieke genetische achtergronden – met o.a. een dominant allel van de *FADING* (*FA*) locus – veroorzaken *ph4* mutaties naast de afwezigheid van vacuolinos in de bloembladepidermis, ook de afbraak van de anthocyaan pigmenten in de centrale vacuole. Dit leidt tot het vervagen (“fading”) van de bloemkleur na opening van de bloemknop. De genetica van dit fenomeen werd al in de jaren 1980 beschreven, maar de onderliggende mechanismen waren nog onbekend. **Hoofdstuk 3** beschrijft de isolatie en karakterisering van het *FADING* (*FA*) gen, de sleutelfactor in het proces van pigmentafbraak. FA codeert voor een lid van de ferro-reductase-oxidases (FROs) uit de familie van metalloreductase-enzymen en komt tot expressie in bloemblaadjes en verschillende vegetatieve weefsels, zoals bladeren en stengels.

In epidermale bloembladcellen van mutanten met een defect MBWW-complex (*an1*, *ph3* en *ph4* mutanten) accumuleert FA op de tonoplast van de centrale vacuole waar de anthocyaan moleculen zijn opgeslagen. Deze cellen missen vacuolinos en FA accumuleert in de centrale vacuole waar het (waarschijnlijk indirect) anthocyanaan destabiliseert. In epidermale cellen van wild type bloemblaadjes, welke vacuolinos
bevatten, accumuleren GFP- en RFP-fusies van het FA eiwit in eerste instantie in vacuolinos, net als andere vacuolaire eiwitten. In de vacuolinos wordt FA-GFP (of ten minste de GFP-tag) uit de membraan vrijgemaakt om vervolgens op te hopen in het lumen van de vacuolinos. Dit zou kunnen verklaren waarom FA-GFP van de vacuolino’s niet verder reist naar de centrale vacuole, zoals andere vacuolaire eiwitten doen, en waarom de FA expressie in wild type bloemblaadjes niet leidt tot afbraak van de anthocyane in de centrale vacuole. In de bloembladepidermis van ph4 en ph3 mutanten ontbreken vacuolinos en kan FA(-GFP) via een directe route de vacuole bereiken, net als in bladcellen, om de afbraak van anthocyane daar te activeren. Deze bevindingen suggereren dat vacuolinos een rol hebben als poortwachter / controlepunt en bepalen dat sommige eiwitten mogen doorreizen naar de centrale vacuole, terwijl andere eiwitten uit de buurt van de centrale vacuole worden gehouden. Deze nieuwe inzichten in de controle van bloempigmentatie (en fruitpigmentatie) via de regulatie van anthocyaan afbraak, kunnen nuttig zijn om gewasvariëteiten te genereren met nieuwe bloemkleuren of verbeterde voedingswaarde van fruit.

Eerder onderzoek toonde aan dat het MBWW-transcriptiefactorcomplex essentieel is voor de vorming van vacuolinos. Waarschijnlijk stuurt MBWW de vorming van vacuolino’s op een indirect wijze, via de activering van ondergeschikte genen coderend voor eiwitten met een directe rol in het verkeer van membraanblaasjes en vorming van vacuolinos. Het mechanisme waardoor vacuolinos worden gevormd en de (MBWW-gereguleerde) genen en eiwitten die erbij betrokken zijn was echter onbekend. **Hoofdstuk 4** presenteert de identificatie van een eerste gen, RAB5a, dat door WMBW wordt gereguleerd en nodig is voor de biogenese van vacuolinos. RAB5 is een lid van de RAB-familie van kleine GTPases, waarvan er vele een cruciale rol spelen bij het specificeren van de ‘identiteit’ van membranen en de
binding van- en herkenning tussen blaasjes en doelmembranen. De identificatie van RAB5a, een homolog van dierlijke RAB5 eiwitten ,toonde aan dat planten-RAB5s evolutionair veel diverser zijn dan tot nu gedacht en bestaan uit drie verschillende fylogenetische groepen, die in petunia worden vertegenwoordigd door RAB5a, RAB5a1 en RAB5a2 plus de homologe (en goed bestudeerde) Arabidopsis eiwitten RHA1 en ARA7. De in hoofdstuk 4 gepresenteerde gegevens tonen aan dat het RAB5a gen wordt geactiveerd door het MBWW-complex en dat het RAB5a-eiwit nodig is om de fusie van een subpopulatie van prevacuolaire compartimenten (PVCs) te bwerken te maken aldus vacuolinos te vormen. Interessant is dat RAB5a1 en RAB5a2 zich in andere cellulare compartimenten bevinden dan RAB5a, en RAB5a ook niet functioneel kunnen vervangen. Dit benadrukt de centrale rol van de RAB5a GTPase bij de vorming van vacuolinos en geeft inzicht in de diversificatie van verschillende vacuolaire compartimenten tijdens de evolutie, door duplicatie en neo-functionalisatie van genen die coderen voor belangrijke regulatoren van het intracellulaire verkeer van membraanblaasjes.

Het verlies van vacuolino's in rab5a mutanten en de vorming van vergrote vacuolinos door expressie van RAB5a vanaf de constitutieve 35S-promoter in Petunia bloemen, gaat gepaard met veranderingen in de afmetingen en vorm van de cellen in de bloemblad epidermis. Een dergelijk effect op epidermale cellen kan de kleur van de bloembladen en zelfs de voorkeur van dierlijke bestuivers veranderen. Een verdere analyse van de effecten van RAB5a-activiteit op de celstructuur kan mogelijk additionele inzichten in de biologische functie(s) van vacuolinos verschaffen.

De zeven bekende PH loci in petunia zijn ontdekt via mutanten met blauw-violette bloemen in plaats van de wilde type rood-violette kleur. Deze PH loci zijn essentieel voor de verzuring van de centrale vacuolen in de epidermis van bloembladen en
daardoor de protonering van de anthocyaan pigmenten en het ontstaan van een rood-violette bloemkleur. Tot dusver waren zes van de zeven PH loci moleculair geïdentificeerd en hun biologische functies beschreven. Het PH7 locus was tot dusver nog niet gekarakteriseerd. **Hoofdstuk 5** beschrijft de moleculaire identificatie van het PH7 gen, middels een vergelijking van de transcriptomen van bloemblaadjes van een wild type lijn (PH7), een instabiele ph7 mutant met een transposon insertie, en een stabiele ph7 mutant. PH7 codeert voor een monosaccharidesuiker transporter die sterk lijkt op – en vermoedelijk homoloog is aan – de vacuolaire glucose/proton symporter ERD6-like6 van Arabidopsis. Mutaties die het PH7 eiwit op verschillende punten afkappen resulteren in duidelijke defecten in de vacuolino-route. Afhankelijk van het punt waar PH7 is afgekapt blokkeert ofwel de vorming van vacuolinos (en mogelijk latere stappen), dan wel het transport van vacuolinos naar de centrale vacuole. Dit laat zien dat PH7 betrokken is bij meerdere stappen in de vacuolino-route en dat PH7 via verschillende mechanismen en domeinen bijdraagt aan de vorming van vacuolino's en het daaropvolgende transport van vacuolinos naar de centrale vacuole.

Deze bevindingen leveren een belangrijke ondersteuning voor eerdere hypotheses dat membraan-transporters additionele functies hebben in membraan verkeer en mogelijk als structurele component fungeren in de herkenning en uiteindelijk fusie van membraanblaasjes en doelwit membranen.
List of Publication


Li S.*, Passeri V.*, Strazzer P., Guzzo F., Spelt C.E., Vandenbussche M., Bliek M., Koes R., and Quattrocchio F.M. FADING, a vacuolar metalloreductase that destabilizes anthocyanins in plant vacuoles (*Co-first authors; in preparation)
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而立未立，惑而不惑。感父母生我养我之恩， 慨外婆牵我挂我之情，谢岳父
母知我信我之义。此恩此情此义，无以为报。唯望早日安生立命，以尽孝道！
芳华易逝，真爱永藏。几多欢乐仅你与我分享，几多艰难只你与我肩扛。未
有你身后默默的付出，哪来我台前的光芒。颗颗，此生遇你，幸也！余生有
你，足矣！只望，执子之手，与子偕老！

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