



# **Tobacco Alkaloid Genetics (TAG) 2017 Report**

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# Objectives

- 1. To understand the genetics that control alkaloid formation in tobacco plants.**
- 2. To understand the feasibility of conventional and non-conventional breeding techniques to modify alkaloid formation in tobacco plants.**
- 3. To understand the impact of tobacco alkaloid levels on leaf production and quality.**

## ❖ Before 2017 CORESTA AP meeting

- To complete the collection of various kinds of publicly available literature and data related to tobacco alkaloids.
- To cluster the alkaloid related factors into different categories including tobacco alkaloid biosynthesis genes, transporters, and regulators.
- To list the known mechanisms of tobacco alkaloid genetic control.

## ❖ Before 2018 CORESTA Congress

- To complete the collection of various kinds of publicly available literature related to tobacco breeding technology.
- To identify conventional and non-conventional technologies used in tobacco breeding to modify alkaloid levels.
- To make a summary of techniques used in modification of tobacco alkaloids.

## ❖ Before 2019 CORESTA AP meeting

- To complete the collection of various kinds of publicly available literature related to the evaluation of tobacco quality and production from tobacco plant containing different alkaloid levels.
- To draft a final report of the TAG TF.

## ❖ Work had finished

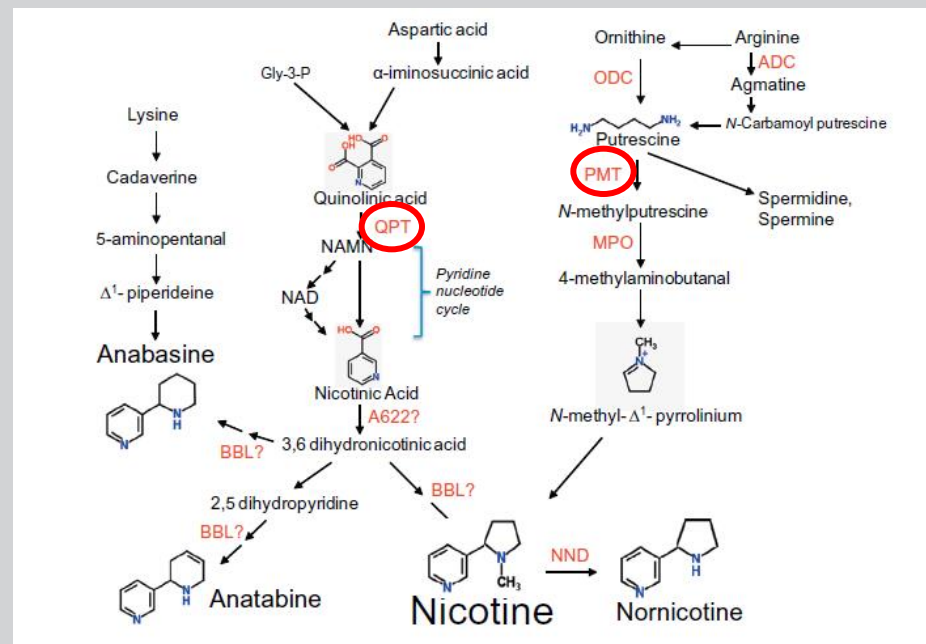
- Collected available literature and data related to tobacco alkaloids
- Alkaloid biosynthesis
- Alkaloid transporters
- Alkaloid regulation



- ❖ Commercial tobacco cultivars: Alkaloid 2%-4% of total dry weight
    - Nicotine ----- 90% of the total alkaloid content
    - Nornicotine
    - Anatabine
    - Anabasine
- } Nearly 10%



- ❖ **Key rate-limiting enzyme**
- ❖ **PMT**: putrescine methyltransferase;
- ❖ **QPT**: quinolinate phosphoribosyltransferase.
- ❖ **ODC**: ornithine decarboxylase;
- ❖ **ADC**: arginine decarboxylase;
- ❖ **MPO**: N-methylputrescine oxidase;
- ❖ **BBL**: berberine bridge enzyme-like;
- ❖ **A622**: isoflavone reductase-like protein;
- ❖ **NND**: nicotine N-demethylase;

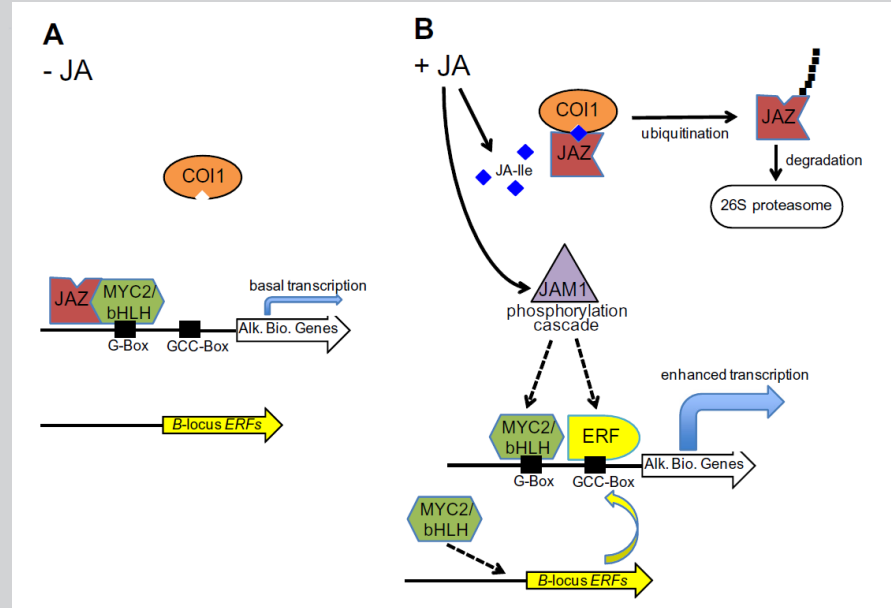


Dewey and Xie (2013)

**Fig. 1. Schematic diagram of alkaloid biosynthesis in *N. tabacum*.**



- ❖ ERF:189, 221
- ❖ MYC2/bHLH
- ❖ COI1,JAZ
- ❖ JAM1

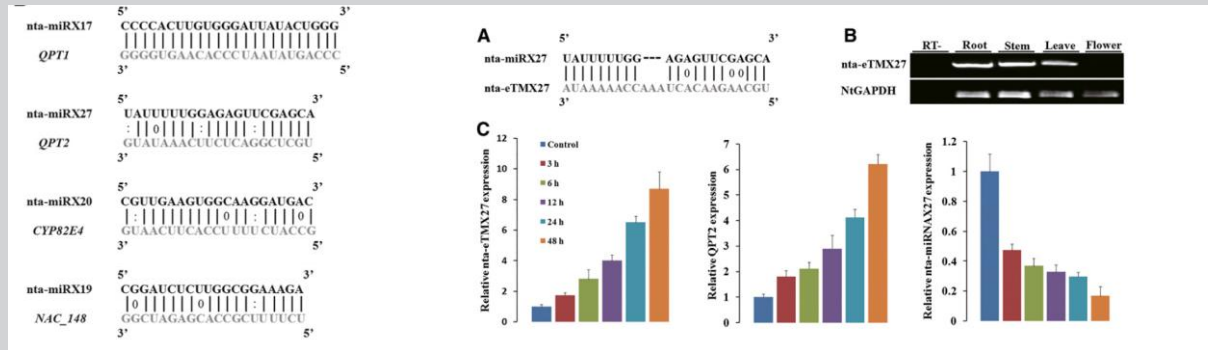


Dewey and Xie (2013)

**Fig. 3. Model depicting our current understanding of JA-stimulated induction of alkaloid biosynthetic genes in tobacco**



- ❖ Nicotine Biosynthesis-Related miRNAs(miRX17,27)
- ❖ Novel tobacco miRNA (nta)-eTMX27 in regulation of nicotine biosynthesis by acting as a decoy of nta-miRX27 to sequester and degrade nta-miRX27 that targets *QPT2*.



Fangfang Li et al. (2015)

**Fig. 4. Topping-induced expression of endogenous nta-eTMX27 in tobacco root**



# Group meeting-Brazil 2017

❖ Meeting Sunday 22 Oct

❖ Acknowledgements:

- Dongmei Xu
- Fabienne Mornet



# THANK YOU